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GENETICS OF TRIATOMINE BUGS
(FAMILY REDUVIIDAE) IN RELATION TO THEIR ROLE AS
VECTORS OF CHAGAS' DISEASE, TRYPANOSOMAL INFECTION OF THE AMERICAS

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A thesis submitted for the degree of

DOCTOR OF PHILOSOPHY

University of London

1976



ACKNOWLEDGEMENTS

I should like to express my gratitude to the Head of the Department of Entomology at the London School of Hygiene and Tropical Medicine, Professor D.S. Bertram, for his continued support and encouragement throughout this research. I thank the technical staff who provided the insect material, Mr. H. Marashi, Mr. R.C. Page and especially Mr. C. Constantinou. I acknowledge the helpful discussions with Dr. J.W. Patterson relating to insect techniques and the advice given on computer programming by Mr. T.F. de C. Marshall, of the Department of Medical Statistics and Epidemiology, London School of Hygiene and Tropical Medicine. I also thank Dr. D.S. Ketteridge and Professor G.H.R. Krombein of the Department of Entomology, London School of Hygiene and Tropical Medicine, for their guidance in techniques with infected bugs and for the use of their equipment. Finally, I acknowledge receipt of a grant from the Overseas Development Ministry, who financed this project.

"The aim of life is self-development. To realize one's nature perfectly - that is what each of us is here for".

Oscar Wilde

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ABSTRACT

Trypanosomiasis caused by Trypanosoma cruzi and transmitted by triatomine bugs is widespread in South and Central America, human infections, chronic or acute, being known as Chagas' disease. Since blood forms of the trypanosome in man and animals are commonly too few to be found in blood films, conclusive evidence of infection is still by xenodiagnosis. This requires vector bugs to be engorged on a suspected infected vertebrate and the bug faeces examined a month or so later by which time the trypanosomes should have greatly multiplied in the insect gut. It is known that there is variation between triatomine species and between individuals within species in ability to develop T. cruzi infections. The present research investigated the genetic basis of this variable susceptibility in one species, Rhodnius prolixus, by selection for susceptible and refractory bug populations and revealed evidence of polygenic control of susceptibility to infection with T. cruzi; this contrasts with the major gene mechanisms reported for other pathogens in other insect vectors. Male bugs showed significantly higher levels of infection than females and may be more efficient for xenodiagnosis.

The radioresistance of triatomine bugs may be related to their chromosome morphology. Male R. prolixus given a sub-sterilizing irradiation dose were less sterile than their progeny, this delay resulting from the diffuse structure of the centromeres of triatomine chromosomes. These results are discussed in relation to possible control of triatomines by genetic manipulation.

Investigation of spermatogenesis by autoradiographic techniques in male R. prolixus revealed that starvation induced diapause inhibits

apoptogenesis. A blood-borne factor produced when the diapausing bug is fed may directly affect the rate of meiosis.

The karyotype is described for several triatomine species. A method was developed for distinguishing between individual mitotic chromosomes within the complements of Triatoma infestans and R. prolixum, the chromosome markers being valuable for further genetic studies of triatomine bugs.

GENERAL INTRODUCTION

Chagas' disease is a serious clinical condition and constitutes one of the most widespread and dangerous human diseases in many South and some Central American countries. This disease is virtually incurable, nor is there an effective prophylactic drug; it can give rise to severe cardiac abnormalities and mega conditions of the alimentary canal and other organs with death from heart failure and other causes. It has been estimated by highly qualified biologists expert committees (W.H.O., 1955, 1956 and 1958) that 40 million people may be exposed to the risk of infection and that at least 7 million people were actually suffering from Chagas' disease. The majority of these cases occurred in Brazil, where it was estimated in 1955 that there were 3 million cases of Chagas' disease (Machado, 1956). Other countries in South America have also reported large numbers of human Chagas' disease: Argentina, 190,000; Colombia, 1,5 million (Garcia, 1951); Ecuador, 80 thousand cases, 1954; Cuba, 75 thousand cases, 1955-1956. Recently the cases have been reported from other South American countries, although the disease is endemic to all of them. Human cases are not normally found beyond northern Mexico (Bustillo 1950 and Argentea 1950) and Chile (Chavez, 1951).

The disease was first described by the Brazilian scientist, Carlos Chagas, who first demonstrated some of a protozoan in the gut of a hemipteran bug, Triatoma dimorpha, in 1905 (Chagas, 1905). These bugs proved to be infective to laboratory animals and Chagas subsequently described the life cycle of Trypanosoma in its development within the host, and the presence of the disease in the Chagas, 1908, 1911). There are also the link to parasite found in a severely sick and Trypanosoma in the blood, the parasite

Dasyatis nasuta (Bragas, 1912), and the first to show that tritosteine
 was the active principle (Bragas, 1912). (Bragas, 1912, p. 108). It
 showed that the trypanosome completed its development in the hind-gut
 of the insect and that the insect was infective (Bragas, 1912) but
 thought that the trypanosomes developed in the salivary glands of the
 bug and were transmitted by the bite. But (Bragas, 1912) did not support
 hypothesis and demonstrated that T. vivax was indeed transmitted in
 the bug faeces.

Thus (Bragas, 1912) has given a complete description of the life cycle
 of T. vivax in the insect host. The life cycle of T. vivax in the
 infected bug faeces first invade cells of the reticulo-endothelial
 system near the site of inoculation. (Bragas, 1912, p. 108). They
 enter the bloodstream and spread to various parts of the body where
 they invade cells of different organs and again multiply. Different
 organs of T. vivax may include all blood organs (heart, liver,
 gut muscle (including heart muscle), spleen, testes, lungs and
 reproductive organs are all favoured sites. On entering a cell the
 body of the trypanosome swells, roundens and its flagellum
 disappears. These developments are followed by heavy division and are
 transformed into trypomastigotes via an intermediate epimastigote
 stage. After 5 days the host cell becomes distended, its cytoplasm
 having been consumed by the parasites, forming a pseudocyst which then
 bursts releasing the trypanosomes into the bloodstream. The trypanosomes
 do not divide in the blood and the numbers present vary with the stage
 of infection; in man they are only numerous during the acute phase of
 the disease and after 4 weeks may be too few to be detected in the
 bloodstream (Bragas, 1912). (Bragas, 1912, p. 108). In the bloodstream
 blood forms are so few as to be seldom or never seen in routine blood
 films. (Bragas, 1912, p. 108). Although the parasites are numerous in the

in the tissues and be released into the blood intermittently.

Dias (loc. cit.) and more recently Brack (1968) have described the life cycle of the trypanosome in the bug, where its entire development takes place in the gut. The length of the cycle in the bug varies according to the stage of development of the host, lasting 5-7 days in young larvae and 10-15 days in adults. A few hours after an infecting bloodmeal, short epimastigotes appear in the flagellum which in the gut of the bug and after 14-20 hours are replaced by amastigotes with a short internal flagellum which later protrudes; the resultant amastigotes along with numerous free trypomastigotes. These then multiply by binary fission and after about 24 hours divide and give rise to small trypomastigotes and epimastigotes. These in turn give rise to blood-meal epimastigotes which appear in the rectum where they attach themselves to the epithelium and become sphaeromastigotes with the flagellum extending into the lumen. These develop into short trypomastigotes which elongate and by the seventh to eighth day slender metacyclic trypanosomes are produced which do not divide and are passed with the faeces of the bug.

Arising from the paucity of trypanosomes in the peripheral blood of the chronically infected vertebrate host - human or other animal (the avian hosts which tritomonins do feed on as well as are susceptible to T. cruzi infection) - and this substantial multiplication of the trypanosomal organism within the bug is the basis of the xenodiagnostic test. This, bug can be reared, or possibly infected, in the laboratory or animal and kept for several weeks until a patent rectal infection is detectable, or not. This is a special aspect of vector function for application to some zoonotic disease systems. It is mentioned genetically in this thesis as the xenodiagnostic test, despite several

conventional tests are applied to the diagnosis of Chagas' disease, which still provides the critical parasitological evidence of infection in man and animal.

Barretto (1968) lists 86 species of American Triatominae, 38 of which have been found to be infectious (14 T. infestans, 10 T. dimidiata, 14 T. sanguinolenta, 10 T. pallidum, 10 T. brevipennis, 10 T. phyllotis, 10 T. lutz, 10 T. rubrocauda, 10 T. melanopus, 10 T. sanguinolenta, 10 T. brevipennis, 10 T. phyllotis, 10 T. lutz, 10 T. rubrocauda, 10 T. melanopus). This occurrence has been associated with variation in the habits of the bug which in endemic areas have become adapted to living in human dwellings and it is these domiciliated rather than sylvatic species which are responsible for most of the transmission of Chagas' disease to man. Barretto (1968) states that the sylvatic species produce very few cases of the disease and that of the domestic species T. infestans and R. prolixus have the widest distribution and are the most important vectors. The principal vectors of Chagas' disease in South America are shown in Table 1.

Table 1. Principal vectors of Chagas' disease in South American countries with high incidence of Chagas' disease.

Country	Species	Reference
Chile	<u>Triatoma infestans</u>	Neiva and Lent (1943)
Argentina	<u>T. infestans</u>	Abalos and Wygodzinsky (1951)
Colombia and Ecuador	<u>Shoenus prolixus</u> <u>T. dimidiata</u>	Dias (1952)
Venezuela	<u>T. sanguinolenta</u> <u>Triatoma prolixus</u>	Dias (1952)
Brazil	<u>T. infestans</u> <u>T. dimidiata</u> <u>T. sanguinolenta</u>	Barretto (1968)

Because of their importance as vectors of Chagau's disease, triatomine bugs have been the subject of a great deal of research work in relation to their distribution, vector potential and control, and one species in particular (Triatoma prolixus), has long been established as laboratory colonies and used for basic researches on insect physiology besides studies bearing on its vector function. However, very little research effort has been applied to the genetics and cytogenetics of triatomine bugs, despite their medical importance, not withstanding that for other medically important insects genetic research has expanded in recent years because of the possibility of controlling vector and pest populations by genetic selection, either at chromosomal or gene level - methods which have the great advantage over conventional pesticide usage of not polluting the environment. The lack of genetic research in triatomine bugs can be related directly to the biology of these insects; genetic experiments inevitably require that successive generations of a given species be reared which, if the work is to be completed within a reasonable time-span, means that organisms with short generation times are preferred.

In the case of insect vectors of disease, the Diptera have become common laboratory tools for genetic research, not only because many of them are pests and vectors, but also because most of them have short generation times and many require relatively simple techniques for colonization. Thus, large amounts of useful information have been gained in relatively short periods of time about, for example, mosquito genetics, and with relatively small budgets.

Triatomine bugs, on the other hand, do not make ideal laboratory tools for genetic research. Their minimum life span from egg to adult, generation time (5-6 months) is the minimum period from egg to adult,

While some species occupy considerably longer times than this) which allows only two generations per year possible at most. Also, because they are obligatorily hemaphysogous, these ticks are expensive to keep in large numbers as they need to be fed and take large blood-meals on their hosts (insects, guinea-pigs, cattle, horses, etc.) at frequent intervals. Raising all or kinds of vectors seems as well as raising of some hosts. This is a labour-intensive process and correspondingly expensive in man-power and vertebrate hosts.

the original material at the U.S.N.I. and T.A.C. provided a large amount of material needed for the present study. Only three T. lewisi (two from Peru and Paraguay) having been kept for many years and hence well adapted to laboratory conditions. The following series were bred in the U.S.N.I. from stock which were long standing laboratory cultures and others built up more recently from bugs supplied from various sources in South America:-

Tetrahymena pallidissima Goshima, 1933. With special note included in Goshima, 1936, 1, by Dr. John Thomas de KUTLA and James HENNINGSEN at L.S.H. and T.Y.

T. latitarsis (Fing. 1914). This species was recorded from Venezuela by Graham Powell and published at B.N. and T.N. in 1951 but the possible origin is unknown. It was subsequently mixed with T. latitarsis received from Argentina, Brazil and Venezuela in 1956.

T. 10111 ²Detritus and detritus, 1971. This species was collected in 1971, found by T. 10111, and also collected at L. 10111 and T. 10111.

T. multicolor (Günther, 1860). This species was brought to the U.S.A. and N.P. by Mrs. H. F. Smith from a collector in

Alaquen, Colombia.

T. tibio-maculata (Pinto, 1969). This species was collected in Santa, Brazil by Dr. H.M. Wilson and colonized at U.S.N. and T.M. in 1973.

T. pallidus (Macneil and Hill, 1961). This species was introduced by Hamilton Macneil, Department of Zoology and Entomology at U.S.N. and T.M. in 1972. Its origin is not known.

T. guatemalensis Hill, 1964. The parental colony and T.M. colony was supplied by Cambridge University in 1961 and originated from L.H.B. and T.M. in the United States. This species was introduced to Santa from Mexico.

T. viridius Hill, 1969. This species was collected in Santa, Brazil, by Dr. H.M. Wilson and colonized at U.S.N. and T.M. in 1973.

Emmetomyia nebulosa (Macneil and Hill, 1961). This species was collected in Santa, Brazil, by Dr. H.M. Wilson and colonized at U.S.N. and T.M. in 1971.

Emmetomyia ruficornis Coquillett, 1964. This species was collected in Santa, Brazil by Dr. H.M. Wilson and colonized at U.S.N. and T.M. in 1972.

E. nebulosa Hill, 1969. The colony of E. nebulosa was established at the U.S.N. and T.M. in 1973 by Dr. A. Hill. This colony was given to him by Professor H. Smith who obtained them from Venezuela at various intervals 1942-1951. This parental colony was subsequently mixed with E. pallidus in 1961. E. nebulosa was introduced from Caracas, Venezuela.

Trichomyia parvicornis (Vignoli and Boudreau, 1964).

T. parvicornis Hill, 1969. This species was sent by Dr. A. Hill to the U.S.N. and T.M. in 1973. This species was introduced from Venezuela, Brazil and brought together with the parental colony to U.S.N. and T.M. in 1973.

some larvae emerging from the eggs and, following the usual 10-12 day interval to second instar larvae, were fed two weeks later as second instar larvae. One week later these had moulted only as third instar larvae, were fed. The larger instar larvae having took about three weeks to moult and the fed as fourth instar larvae, and so on further down again after another week or two to fifth instar larvae. The final moult to adult occurred after 3 or 4 weeks and adults were fed a week after emergence. Larvae were reared on a 12 hr. feeding interval and after three feeds they were disposed of to make way for more rearing adults. While only a small number of adults of each generation were kept for study, the low cost of rearing, and the large numbers of bugs could be reared without adverse effect on the colonies. This feeding system enabled feeding schedules and rearing work to be planned well in advance.

The time for rearing bugs in this laboratory varied between 10 and 15 days for immature stages and eggs for both species, as already mentioned, with species, the variable stages especially for *E. pallidus*, in which most of the ground work was completed. Other species, such as *Penstemonius marginatus*, could take as long as 9 months or more to complete the life-cycle, *E. pallidus* having a relatively short life-cycle, in the typical stages, of about 3 months. For species other than *E. pallidus*, material was supplied at appropriate stages and condition by other workers in the department from the colonies which they were particularly concerned with rearing for other researches.

Because of its relatively short life-cycle, the ground research was concentrated on *E. pallidus*. From selected ground research, deemed feasible in three years, were chosen for this study as likely to be of particular interest to problems of tristamine systematics, while some other ground research was left to other workers, or to be

of these processes in the expression of numerous biological functions. Some researches are presented in the thesis in four parts, as briefly indicated below.

PART TWO - Cytotaxonomy of the Tristoniinae

The cytotaxonomy of insects is now a well established technique, and can provide great insight into the relationships of different species on the basis of chromosomal differences. The cytotaxonomy or tristoniine bugs has, however, been extremely neglected because these bugs do not possess the giant polytene chromosomes so useful to taxonomists in the diptera. The present work consisted of comparing the karyotypes of the tristoniine species and hybrid species available and developing methods of producing chromosome markers for this group of insects using recently developed fluorescent staining methods.

PART TWO - The regulation of spermatogenesis

Hormones have been shown to control many aspects of growth and differentiation in insect development, but the control factors involved in development of the testis, and the regulation of spermatogenesis in the fully developed testis, remain subject to debate. Most work carried out on the development of insect testes has concentrated on the growth of the testis as a whole and the proliferation of spermatogonia. The present experiments examined the effects of starvation and feeding on the intra-meiotic events of spermatogenesis in M. prolixa fifth instar larvae and adults.

PART THREE - The effect of temperature on the development of the testis

Introduction

The development of the testis is controlled by a number of factors, including temperature. The present experiments examined the effect of temperature on the development of the testis in M. prolixa fifth instar larvae and adults. The results of these experiments are given in the following chapters.

is susceptibility to the parasite infection affecting the outcome. The present experiment was designed to study whether the susceptibility of R. prolixus to T. cruzi is genetically determined by following a selection program to breed refractory and susceptible populations of bugs. Should susceptibility be related to genotype, then it should be possible to breed a highly susceptible strain to improve the efficiency of the transmission test.

PART IX - The inheritance of radiation induced semi-sterility in Rhodnius prolixus

During the past several years, in several ways, we have worked on insect vectors and pests by the release of artificially sterilized males and females. Insect control is a major problem in the tropics. It has been shown that, like the lepidoptera, triatomine bugs are very resistant to sterilization by irradiation, such that doses required for complete sterility of males result in them being unable to compete with normal males in mating. The present experiments assessed the effect of sub-sterilizing doses of irradiation on the fertility of irradiated males, and successive generations of their offspring, and related the changes in fertility to chromosomal damage found in spermatocytes of experimental males. The release of semi-sterile lepidoptera has been used as a means of controlling this radio-resistant group. The purpose of this aspect of the thesis research on triatomines was to assess the fertility of male R. prolixus treated with sub-sterilizing doses of irradiation and to relate these effects to chromosomal changes; the feasibility of controlling this vector species by releasing semi-sterile males would then be properly assessed.

PART ONE

CYTOTAXONOMY OF THE TERLATORIDAE

INTRODUCTION

Insects of the sub-family Triastorinae (Family Reduviidae, Sub-Order Heteroptera, Order Hemiptera) are widely distributed in the Americas and comprise 18 genera, and more than 50 species (Usinger et al., 1961). A great deal of present-day morphological research has been done on the members of this group of insects, and more or less the same also involved experimental hybridization of different species to determine the nature of their relationship from their hybrid fertility or sterility. Morphological experiments have been carried out by Hughes (1961) and comparisons made by de Meillon (1961) on *Triastor* and *Phyllot* and Usinger (1951) on *Triastor*, *Triastor* and *Phyllot* problems in comparison made on a taxonomic laboratory in Germany. Light gray (very small) morphological studies of the Triastorinae, Brodie and Ryckman (1967) examined the antigenic relationships of 18 populations of bugs comprising five *Triastor* species from both North and South America using anti-*Triastor* and *Phyllot*.

Differences in chromosome morphology between species have provided taxonomists in recent years with additional criteria to aid classification. Thus, chromosome characteristics such as number, shape, size and general appearance are all important in the classification of insect species. The cytogenetic karyotype of some potentially important species has indeed been well established (Baker, 1968a). Compared with other disease vectors, however, the cytogenomy of the tabanids are less generally available. Thus, little is known of the karyotype of *Chrysops* (1968b), *Chrysops* (1968c), *Chrysops* (1968d), *Chrysops* (1968e) and *Chrysops* (1968f). There have been suggested studies of the number of autosome and sex-chromosomes for several species of tabanids.

The Triatominae have large numbers of relatively small and almost

indistinguishable autosomes, with usually 20 autosomes plus an XY system in the male, although some species have multiple X-chromosome systems (e.g., *Phlebotomus* and *Simulium*). The only detailed study of the cytotaxonomy of these pests is given by Ueshima (1961, cit.) who describes details of meiotic chromosome behaviour for 20 species and 1 species hybrid from studies of testis squashes. These studies showed, unfortunately, that the meiotic chromosomes of tritoxine spermatocyte nuclei do not provide the taxonomist with a useful cytogenetic tool, because of the similarity of the small chromosomes. A further limitation was that relative arm length of chromosomes, which can be a useful diagnostic feature in many insects, has no systematic value in hemipterous insects which have isochromosomes (e.g., *Phlebotomus*).

Polytene chromosomes, in which minute differences in morphology can be detected, have facilitated the identification of species of Dipteran chromosomes so that sibling species may be distinguished. Such studies of polytene chromosomes in mosquitoes (Kitzinger et al., 1967) black-flies (Dunbar, 1966) and Tsetse-flies (Southern and Pell, 1973) have brought great precision to studies of systematic relationships within these vector species. Polytene chromosomes have also been recognized from any tissue of tritoxine bugs at any developmental stage so that, again, this tool, so valuable in these several vector insects is not available for bug taxonomists.

Cytogenetic studies of *Phlebotomus* and *Simulium* have been handicapped by the nature of their chromosome morphology, and this was also the case for other animal groups, notably for mammals and particularly for human cytogeneticists faced with large numbers of very similar sized chromosomes in the human cell (for a recent review see

Ford, 1973). However, in recent years, the study of mammalian cytogenetics has been revolutionized, Caspersson et al. (1968, 1969a and b) showing that by using fluorescent dyes they could distinguish brighter and darker zones in metaphase chromosomes of man when the preparations were examined under U.V. light on a fluorescence microscope. Later, with the dye quinacrine mustard (Q.M.), Caspersson et al. (1970a, b and c) were able to delineate many chromosome of the human karyotype. The precise nature of these Q-bands, however, remained controversial; it was assumed initially that the alkylating agent, mustard, bound preferentially to regions of DNA rich in guanine and that the fluorochrome quinacrine attached to the mustard would fluoresce most strongly in these regions (Caspersson, 1968). However, it has since been shown that Q.M. has an affinity for adenine-thymine rich DNA regions and not guanine-cytosine regions, which makes the original hypothesis untenable (Weinblum and DeHesseth, 1972). Despite the controversy which has surrounded the mechanism of chromosome fluorescence, its usefulness for delineating different chromosomes was an important advance, although the technique required complex and expensive equipment and the fluorescence of the preparations faded with time. These problems were circumvented when a technique was developed which would allow the production of human chromosomes using Giemsa stain following a pre-treatment incubation in a solution of quinacrine mustard (1968) (Weinblum et al., 1971). Using this Giemsa (G-band) technique higher resolution of bands could be obtained if the preparation was also given a brief pre-treatment in hypoxanthine (Weinblum, 1972). As with the Q-bands, the cytochemistry of G-band production remains a matter for debate; it was thought originally that G-bands reflected a denaturation-reassociation process of repetitive DNA (Sumner et al., 1971) but when the bands were

produced by an enzyme which did not act by denaturation (Gonbright, 1972) this theory was rejected. Kato and Moriwaki (1972) found that bands could be induced in mammalian chromosomes by a variety of chemicals but could suggest no common mode of action. Sanchez and Yunis (1974) have recently suggested that the basic organization of DNA in eukaryotes and, in particular, its repetitive nature are responsible for band production but an ultrastructural study of G-bands (Burkholder, 1975) has shed little light on their nature. Whatever their origin, G-bands have improved the study of mammalian cytogenetics and since the chromosomes of tritamine bugs presented problems akin to mammalian chromosomes (i.e. large numbers of small, karyologically-bivalent chromosomes) it was decided as part of the present work to attempt to apply G-banding techniques to mitotic cells of tritamine material. Although little work has been previously reported on banding of insect chromosomes, several workers have reported successful application of the C-banding technique to insect material. The C-banding technique involves pre-treatment of cells in an alkaline or acid solution followed by S.S.C. incubation and Giemsa staining, which stains a single band of heterochromatin in the centromeric region. Hau (1971) first showed that C-bands could be produced with insect material using carbazole, benzidine, and collidine etc. al., (1972) also successfully stained the supernumerary B-chromosomes of the grasshopper Myrmica doloticus parvulus, using the same technique. Newton et al., (1974) were able to distinguish the X and Y-chromosomes of Perla marginata in larval brain cells by C-banding and Brets and Stoll (1974) demonstrated C-bands in mitotic and meiotic chromosomes of the cricket Gryllus argenteus. However, C-banding provides only a limited system of chromosomal markers and it was hoped that by applying the C-banding technique to insect chromosomes,

a more sophisticated system of insect rearing is required.

In addition, the karyotypes of 13 species of tristamine and one species hybrid were recorded at adult midlife. *Phaenix melania*, *Gillnetta* unreported in tristamine bugs, was investigated in one species, *R. paulownia*.

I now turn to describe the techniques used, for conventional orcein staining of chromosomes and those developed on the principles of G-banding for human chromosomes, for tristamine species and the results obtained regarding the karyotypes for the following tristamine material: - *Tristamine* *luculentus*, *T. luculentus*, *T. luculentus*, *T. luculentus*, *T. luculentus*, *T. luculentus*, *T. luculentus*, *T. luculentus*, *T. luculentus*, *T. luculentus*, *T. luculentus*, *T. luculentus*, and *T. luculentus* x *T. luculentus*.

Materials

1.1.1. *Tristamine*

Tristamine *luculentus*. Tristamine were dissected from 1st adult instar under lowest yellow light (1 g 1000). It is 100% pure, 100% pure, 100% pure, fixed in 3:1 absolute ethanol:acetic acid and stored at 4°C in labelled glass vials. Tristamine were dissected from 1st instar by taking out fixed follicles out on clean microscope slides in lacto-propionic-orcein using a small brass rod with a flat end. The orcein stain was made by combining 1 g pyridine, 100 ml 100% ethanol, 100 ml 100% ethanol, 50 ml 100% propionic acid and 50 ml 45% lactic acid for 24 hours and then filtering. Preparations were squashed by finger pressure under 22 x 22 mm coverslips and sealed with rubber solution to prevent evaporation. After examination and photography, coverslips were removed by freezing in liquid nitrogen and prizing off with a razor

slides, the preparations then being placed into standard slides and mounted in coverslip-mounted mounts. Photographs of the resultant chromosome configurations were taken with Kodacord 35 mm micro-film (Kodak, London, England) developed in FF contrast developer (Ilford, Essex, England) and fixed in Arstix (May and Baker, Dagenham, England).

Figure 1. Diagrammatic illustration of the four panels of all the types of mutation. The polymorphic marker is located on a heterozygous locus representing two chromosomes, one with allele 1, 2 and 3; chromosome 1.

The autosomes do not stain as deeply as the sex-chromosomes in early prophase but at diakinesis they have become hypertensive, and the autosomes can be seen to be joined by a single terminal chiasma in each bivalent.

Autosomes 1-10, 12-13, 15-16, 18-19, 21-22, 24-25, 27-28, 30-31, 33-34, 36-37, 39-40, 42-43, 45-46, 48-49, 51-52, 54-55, 57-58, 60-61, 63-64, 66-67, 69-70, 72-73, 75-76, 78-79, 81-82, 84-85, 87-88, 90-91, 93-94, 96-97, 99-100, 102-103, 105-106, 108-109, 111-112, 114-115, 117-118, 120-121, 123-124, 126-127, 129-130, 132-133, 135-136, 138-139, 141-142, 144-145, 147-148, 150-151, 153-154, 156-157, 159-160, 162-163, 165-166, 168-169, 171-172, 174-175, 177-178, 180-181, 183-184, 186-187, 189-190, 192-193, 195-196, 198-199, 201-202, 204-205, 207-208, 210-211, 213-214, 216-217, 219-220, 222-223, 225-226, 228-229, 231-232, 234-235, 237-238, 240-241, 243-244, 246-247, 249-250, 252-253, 255-256, 258-259, 261-262, 264-265, 267-268, 270-271, 273-274, 276-277, 279-280, 282-283, 285-286, 288-289, 291-292, 294-295, 297-298, 300-301, 303-304, 306-307, 309-310, 312-313, 315-316, 318-319, 321-322, 324-325, 327-328, 330-331, 333-334, 336-337, 339-340, 342-343, 345-346, 348-349, 351-352, 354-355, 357-358, 360-361, 363-364, 366-367, 369-370, 372-373, 375-376, 378-379, 381-382, 384-385, 387-388, 390-391, 393-394, 396-397, 399-400, 402-403, 405-406, 408-409, 411-412, 414-415, 417-418, 420-421, 423-424, 426-427, 429-430, 432-433, 435-436, 438-439, 441-442, 444-445, 447-448, 450-451, 453-454, 456-457, 459-460, 462-463, 465-466, 468-469, 471-472, 474-475, 477-478, 480-481, 483-484, 486-487, 489-490, 492-493, 495-496, 498-499, 501-502, 504-505, 507-508, 510-511, 513-514, 516-517, 519-520, 522-523, 525-526, 528-529, 531-532, 534-535, 537-538, 540-541, 543-544, 546-547, 549-550, 552-553, 555-556, 558-559, 561-562, 564-565, 567-568, 570-571, 573-574, 576-577, 579-580, 582-583, 585-586, 588-589, 591-592, 594-595, 597-598, 599-600, 602-603, 605-606, 608-609, 611-612, 614-615, 617-618, 620-621, 623-624, 626-627, 629-630, 632-633, 635-636, 638-639, 641-642, 644-645, 647-648, 650-651, 653-654, 656-657, 659-660, 662-663, 665-666, 668-669, 671-672, 674-675, 677-678, 680-681, 683-684, 686-687, 689-690, 692-693, 695-696, 698-699, 701-702, 704-705, 707-708, 710-711, 713-714, 716-717, 719-720, 722-723, 725-726, 728-729, 731-732, 734-735, 737-738, 740-741, 743-744, 746-747, 749-750, 752-753, 755-756, 758-759, 761-762, 764-765, 767-768, 770-771, 773-774, 776-777, 779-780, 782-783, 785-786, 788-789, 791-792, 794-795, 797-798, 799-800, 802-803, 805-806, 808-809, 811-812, 814-815, 817-818, 820-821, 823-824, 826-827, 829-830, 832-833, 835-836, 838-839, 841-842, 844-845, 847-848, 850-851, 853-854, 856-857, 859-860, 862-863, 865-866, 868-869, 871-872, 874-875, 877-878, 880-881, 883-884, 886-887, 889-890, 892-893, 895-896, 898-899, 901-902, 904-905, 907-908, 910-911, 913-914, 916-917, 919-920, 922-923, 925-926, 928-929, 931-932, 934-935, 937-938, 940-941, 943-944, 946-947, 949-950, 952-953, 955-956, 958-959, 961-962, 964-965, 967-968, 970-971, 973-974, 976-977, 979-980, 982-983, 985-986, 988-989, 991-992, 994-995, 997-998, 999-1000, 1002-1003, 1005-1006, 1008-1009, 1011-1012, 1014-1015, 1017-1018, 1020-1021, 1023-1024, 1026-1027, 1029-1030, 1032-1033, 1035-1036, 1038-1039, 1041-1042, 1044-1045, 1047-1048, 1050-1051, 1053-1054, 1056-1057, 1059-1060, 1062-1063, 1065-1066, 1068-1069, 1071-1072, 1074-1075, 1077-1078, 1080-1081, 1083-1084, 1086-1087, 1089-1090, 1092-1093, 1095-1096, 1098-1099, 1101-1102, 1104-1105, 1107-1108, 1110-1111, 1113-1114, 1116-1117, 1119-1120, 1122-1123, 1125-1126, 1128-1129, 1131-1132, 1134-1135, 1137-1138, 1140-1141, 1143-1144, 1146-1147, 1149-1150, 1152-1153, 1155-1156, 1158-1159, 1161-1162, 1164-1165, 1167-1168, 1170-1171, 1173-1174, 1176-1177, 1179-1180, 1182-1183, 1185-1186, 1188-1189, 1191-1192, 1194-1195, 1197-1198, 1199-1200, 1202-1203, 1205-1206, 1208-1209, 1211-1212, 1214-1215, 1217-1218, 1220-1221, 1223-1224, 1226-1227, 1229-1230, 1232-1233, 1235-1236, 1238-1239, 1241-1242, 1244-1245, 1247-1248, 1250-1251, 1253-1254, 1256-1257, 1259-1260, 1262-1263, 1265-1266, 1268-1269, 1271-1272, 1274-1275, 1277-1278, 1280-1281, 1283-1284, 1286-1287, 1289-1290, 1292-1293, 1295-1296, 1298-1299, 1301-1302, 1304-1305, 1307-1308, 1310-1311, 1313-1314, 1316-13

Plate 1.1 shows a spermatocyte in diplotene from T. pallidus, which is protracted to the maximum translocational configuration with the four chromosomes associated together, and at this stage becomes negatively heteropycnotic relative to the autosomes and remain so throughout the rest of meiosis; Plate 1.2 shows a metaphase I cell from T. pallidus. The chromosomes are arranged like the 10 in Plate 1.3 at the centre of a ring of autosomes and this arrangement is also illustrated by Plate 1.3 with a metaphase I cell from T. lefeevrei. The karyotypes of the species T. pseudomaculata and T. florida will

be dealt with later in the section, related to the cytogenetics of the hybrids produced by crossing these two species.

Triatomia tibio-aurulata, T. phyllonora, T. protracta, T. lenti and T. vittiger. These 5 species of the genus Triatomia all have multiple non-chiasmate X_nY systems in the male, 4 of them with X_1X_2 and T. vittiger with an $X_1X_2X_3Y$ system. The course of meiosis in these species at the first and the 2nd metaphase is described above. The 20 autosomes arrange themselves as 10 bivalents on the metaphase plate and the sex-chromosomes are usually aligned peripherally. Plates 1.5, 2.6 and 1.7 show metaphase I spreads from T. tibio-aurulata, T. phyllonora and T. protracta and Plate 1.9 shows a full 2d metaphase from T. lenti. At second metaphase in this group of species the sex-chromosomes always lie in the centre of the ring of autosomes and assume the characteristic "heart" and "sp" pairing with 10 autosomes for T. phyllonora in Plate 1.8. At second anaphase the Y-chromosome goes to one pole and the multiple X-chromosomes go to the other pole. T. vittiger is of special interest being the only species examined with three X-chromosomes. Plate 1.10 shows a metaphase I cell of this species, and it can be seen that the Y-chromosome is much larger than all the X-chromosomes being about the size of the autosomes.

Panethosyllus marginatus. This species is characterized by having only 18 autosomes which form 9 bivalents at metaphase I. The sex-chromosomes (X_1X_2Y) lie at the periphery of the spindle at metaphase I (Plate 1.11) and divide independently at the first division. Plate 1.12 shows an anaphase I division. At metaphase II (Plate 1.13) the sex-chromosomes again lie at the periphery of a ring of autosomes.

Phodinus nebulosus and P. pusillus. The two species studied of the genus Phodinus both have 20 autosomes plus an XY sex-system. Their meiotic chromosome number is 2n=21 smaller than those found in the genera



Plate 1.1. Spermatocyte of *E. bispinosus*. $2n = 31 + XY$.

Note sex-chromosomes isopycnotic with autosomes. Ten autosomal bivalents with a single terminal chiasma. X 2500.

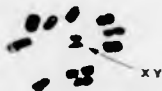


Plate 1.2. Spermatocyte of *J. medusta*. $2n = 20 + XY$.

Note negative heteropycnosis of sex-chromosomes which lie at centre of ring of autosomal bivalents. X 2500.



Slide 14 Metaphase I T. tritici 20 + XY.
X small.



Slide 15 Metaphase II spermatocyte of T. tritici. Note sex
pseudo-bivalent (sb) L. centromere of ring of autosomes.
X small.



Plate 1.5. Metaphase I spermatocyte of *T. tibio-maculata*, $2n = 20 + X_1X_2Y$.
X 2500.



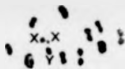
Plate 1.6. Metaphase I spermatocyte of *T. phyllosoma*, $2n = 20 + X_1X_2Y$.
X 2500.



Plate 1.7. Early metaphase I spermatocyte of T. penetrans. $2n = 20 + X_1X_2Y$.
X 2500.



Plate 1.8. Diakinesis spermatocyte of T. lenti $2n = 20 + X_1X_2Y$.
Note very small X_2 . X 2500.



Slide 141. Spermatocyte of *T. castaneum*, $2n = 20 = 8A, 2X, Y$.
Note sex-chromosomes at centre of ring of autosomes and
their characteristic 'touch and go' pairing. X 2500.



Slide 142. Metaphase I spermatocyte of *T. vittiger*, $2n = 20 = 8A, 2X, Y$.
Note large Y-chromosome similar in size to autosomes. X 2500.



Plate 1.11. Metaphase I spermatocyte of P. megistus, $2n = 18 + X_1X_2Y$.
Note only 9 bivalents. X 2500.



Plate 1.12. Anaphase I spermatocyte in P. megistus. X 2500.



Plate 1.11. Metaphase I spermatocyte of P. megistus. X 2500.



Plate 1.14. Metaphase I spermatocyte of R. neglectus, $2n = 20 + XY$.
Note small size of chromosomes compared to Tristoma
species. X 2500.



Plate 1.15. Metaphase II spermatocyte of R. neglectus. X 2500

Trioxys and Panictorhynchus, as seen in Plate 1.14 which shows a metaphase I cell of R. neglectus with the X and Y-chromosomes at the periphery of the spindle. Plate 1.15 shows a metaphase II cell of this species with the XY pseudo-bivalent at the centre of a ring of autosomes. The karyotypes of these 2 Rhombus species are barely distinguishable in meiotic preparations except that the chromosomes of R. punctatus are slightly larger (see later and Plate 2.6, Part Two).

(b) Female Melosis

Cytogenetic technique. The method of Perry and Jones (1951) developed for the study of grasshopper oocyte nuclei was used to study female meiosis in tritosome eggs. Eggs were placed in 311 absolute alcohol, then in cold fixative, then removed with hexane and fixed for 5-10 minutes. The contents of the egg were then squeezed out of the case with forceps, placed in a drop of lacto-propionic-orcein on a siliconized coverslip ('Repakote', Hopkin and Williams, Essex, England) and allowed to stain for one hour. The coverslips were then inverted on to clean slides, squashed and examined for the egg nucleus.

Results. Because of the large numbers of cells in division in a testis these organs have always been favoured for the study of meiosis. The female reproductive system, however, presents considerable technical difficulties in demonstrating meiosis. In insects it is complicated by large amounts of yolk and therefore presents considerable technical difficulties in demonstrating meiosis. (Cytogenetic technique). Perry and Jones (loc. cit.), however, developed an oocyte squash technique which proved to be suitable for application to tritosome eggs. Since a large colony of R. prolixus was being maintained for the present work the opportunity existed to examine large numbers of eggs of this species. It was found that the time of oocyte division of the ovum to occur at the time of ovulation following the

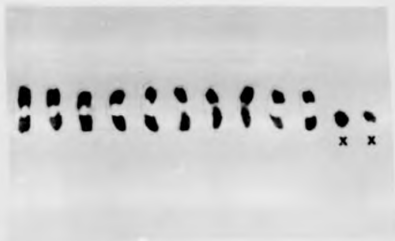


Plate 1.16. Metaphase I oocyte nucleus from an egg of E. prolixus.

Cut-out karyotype shows $2n = 20 + XX$. X 2500.

entry of the sperm into the plasma of the egg and for fertilization of the ovum nucleus to take place just before the egg is laid (Wigglesworth, 1967). The female reproductive organs of B. prolixus consist of 2 ovaries each made up of several ovarioles of the telotrophic type in which the nurse cells are confined to the apex of each ovariole and are connected to the developing oocytes by long tubular oocyte chorion (Wigglesworth, 1967; Fagan, 1967). Oogenesis occurs when the leading oocyte of each ovariole is ripe and the eggs pass one after the other into the lateral, on their way to the common, oviduct. Examination of large numbers of eggs from the reproductive tracts of female B. prolixus at all stages of development up to the substantial yolk laden egg at oviposition revealed the oocyte nuclei, but this may have been due in part to the technical difficulties of locating the small nuclei in the mass of yolk in the eggs. The 3 oocyte nuclei which were found at first metaphase were all located in the lateral oviduct and therefore close to the oviduct. One of these female metaphase I nuclei is shown in Plate 1.16 which shows that the female has 10 bivalents of very similar size with 2 X-chromosomes.

(c) G-Banding of Mitotic Chromosomes

Published photographs of mitotic chromosomes of triatomine bugs have shown groups of chromosomes that I & B banded chromosomes (Wigglesworth, 1967; Wigglesworth, 1967; Wigglesworth, 1967). Wigglesworth (1967) examined mitotic cells in the fat body, epidermis and muscle of B. prolixus larvae and his results also showed minute mitotic metaphase chromosomes. Examination of these tissues in the course of the present work confirmed these observations - mitoses in these asexual tissues appeared to be similar to those found in gonadal mitoses (Part Two, Plate 2.1). However, studies by Mellanby (1936a and b) of the embryonic

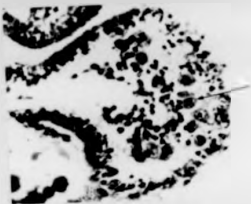


Plate 1.17. 1.5. of embryo of *Trichostema* 5-days old showing large mesodermic vesicles. X 800.

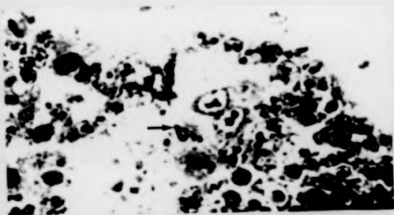


Plate 1.18. 1.5. of embryo of *Trichostema* showing large mitotic metaphase chromosomes of mesoblasts. X 800.

development of the embryo contained special large, neuroblast-like cells present in the embryo at the time of head and thoracic segmentation. Mellanby (1936a) found that these large cells were neuroblasts responsible for the formation of the ventral nerve cord; they appeared at day 5 of embryonic development. Sections of 5-7 day-old eggs were, therefore, studied (using 1. infertile eggs initially) and large neuroblasts resembling those reported by Mellanby (1936a) for the embryo were found. Sections of eggs were prepared using the method described by Anderson (1941) for ferret eggs, and were stained with hematoxylin and eosin. Plates 1.17 and 1.18 show sections through 1. infertile eggs and demonstrate the neuroblast cells.

Cytogenetic technique for G-banding mitotic chromosomes. Squash preparations of 5-7 day-old 1. infertile eggs confirmed that the chromosomes of these neuroblasts were indeed larger than those found elsewhere in the embryo, hence in other tissues. Having located these cells with large chromosomes which seemed suitable for G-banding the next technical problem was to produce a suspension of these cells. It is an essential pre-requisite for the G-banding technique to have a suspension of living cells which, when dropped on to a slide and collected, produces a preparation with individual cells randomly dispersed over the slide and showing good metaphase spreads. Cytogeneticists working with human material routinely employ a technique developed by Moorhead et al. (1960) for lymphocyte cultures to produce colonies.

Five to seven day-old embryos were dissected from the triatomine eggs in L15 medium (Leibovitz, 1961) and placed in tyrocyte (1960) citrate for 8 minutes. The embryonic cells were then dispersed in 0.25% trypsin (Boehringer-Mann, Gibco, Grand Island, N.Y.) for 10 to 15 minutes.

versene (0.1 g EDTA, 4.0 g NaCl, 1.1 g KCl, 0.58 g Na_2HPO_4 , 0.1 g CaCl_2 , and 0.1 g glucose in 500 ml distilled water), the resulting cell suspension washed in insect saline and centrifuged at 1000 r.p.m. for 5 minutes. The supernatant was discarded and the pellet flicked to form a thin layer up the walls of the centrifuge tube which was rapidly fixed by adding drops of freshly made 3% absolute ethanol:acetic acid fixative on the cells. After standing for 5 minutes the cells were centrifuged at the same speed, re-suspended in fresh fixative and, after standing for a further 10 minutes, were centrifuged and finally suspended in a few drops of fixative. Following penetration and wash by the method of Evans et al. (1964), the suspension being dropped on to acid-cleaned slides and blown dry when the interference rings appeared. The preparations were kept for one week in dust-proof boxes before further treatment, since attempts to produce band preparations immediately after fixation were unsuccessful. The G-banding technique employed was essentially that described by Sumner et al. (1971); slides were incubated for one hour in 0.1 M citric acid, sodium chloride 0.05M, 0.05M trisodium citrate at 60°C, rinsed in absolute ethanol and stained in 0.5% Giemsa stain in 0.1M citric acid, sodium chloride 0.05M, 0.05M trisodium citrate at pH 6.8 for 3 minutes. Fresh buffer was prepared by diluting 1.5 ml of 0.1M citric acid to 50 ml with distilled water, and adjusting the pH to 6.8 with 0.1M sodium phosphates. After staining, the preparations were rapidly rinsed by running distilled water into the Coplin jars and finally removing to be air-dried and mounted in Euparal.

The results of using this technique with neuroblasts of 2 species of Chironomus and Simulium are shown in Figs 1, 2 and 3.



Plate 1.12. Cut-out karyotype of *E. prolepis* from neuroblast of male embryo G-banded by A.S.G. technique.



Plate 1.20. Cut-out of karyotype of *T. infestans* from neuroblast of male embryo G-banded by A.S.G. technique.

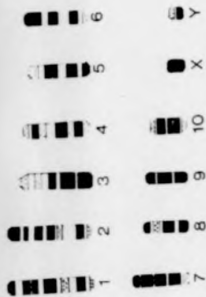


Plate 1.21. Idiogram of mitotic chromosome complement of *R. prolixus*.



Figure 1.111 Human karyotype showing normal complement of 46 chromosomes.

It is clear from a study of these photographs that very definite distinctions can be made between chromosome pairs within complements and that G-bands provide excellent markers of particular chromosomes. Comparison of the 2 out-of karyotypes shows a great deal of interspecific similarity which is expected since both species belong to the same subgenus. The 2 idiograms prepared from these karyotypes (Figures 1.12 and 1.13) provided a means of identifying chromosomes within complements. Comparison of these idiograms revealed sufficient distinguishing features for the recognition of each pair of homologues within each species and although the bands are not as detailed as those found in the large Dipteran genomes, these results are noteworthy as representing an advance in triatomine cytogenetics which has so far been limited to the study of male meiotic chromosomes conventionally prepared and stained with orcin as illustrated by the numerous photographs (Plates 1.1 - 1.15) of meiotic karyotypes in the early sections of this study.

(d) Hybrid species males

Dz. A. Perlowagon-Szumlewiec has been studying the fertility of F_1 hybrids formed by crossing various species of triatomine bugs with a view to the production of sterile males which could perhaps be used in a programme designed to control triatomine populations by releasing sterile males. He has found that the cross *T. parvulus* ♀ X *T. lundii* ♂ and its reciprocal cross were both semi-fertile and the F_1 males from these crosses were always completely sterile, while the F_1 females had very low fertility. For this reason, the F_1 males were selected for cytogenetic study. The F_1 male sterility was cytogenetic in origin and also some F_1 and parental males of both species to London for cytogenetic analysis, the results of which are presented here.

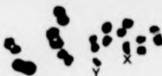


Plate 1.23. Metaphase I spermatocyte of T. pseudo-maculata, $2n = 20 + XY$.
X 2500.

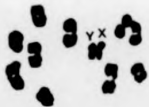


Plate 1.24. Metaphase I spermatocyte of T. sordida, $2n = 20 + XY$.
X 2500.



Plate 1.25. Metaphase I spermatocyte of *T. pseudo-maculata* ♀ X *T. mordax* ♂
F₁ hybrid. Note chain of III and univalents. X 2500.



Plate 1.26. Metaphase I spermatocyte of *T. mordax* ♀ X *T. pseudo-maculata* ♂
F₁ hybrid. Note chain of VII and ring of V associations plus
univalents. X 2500.

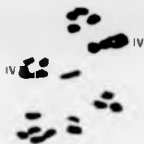


Plate 1a, 1b. Metaphase I spermatocytes of *T. arida* x *T. pseudo-nuculita* δ
 F_1 hybrid. Note chain of IV and ring of IV associations.
 X 2500.



Plate 1a, 1b. Metaphase I spermatocytes of *T. arida* x *T. pseudo-nuculita* δ
 F_1 hybrid. Note chain of IV association and 10 univalents.
 X 2500.



Figure 14.10. Diskineate spores of *S. longicauda* X *T. longicauda*
 F_1 hybrid, 8 bicolors and 8 unicolors. X 2500.

Parental strains

(i) T. pseudo-maculata. The karyotype of this species shows no differences at male meiosis from that of T. maculata. Plate 1.23 shows a metaphase I cell from T. pseudo-maculata with 20 autosomes plus an X-Y system.

(ii) T. sordida. This species also has 20 autosomes and an XY system although the chromosomes of this species are all somewhat larger than those of T. pseudo-maculata (Plate 1.24).

F₁ hybrids

Meiosis in the F₁ hybrid males formed by crossing these 2 species was characterized by the formation of multiple associations and univalents. Multiple associations assumed a variety of forms in the hybrid males, the most common being chains which varied in size from XIII (Plate 1.25) to chains of VII (Plate 1.26). Although chains were the most frequently seen in multivalent associations, ring multivalents were also found in these hybrid males and they too varied in number of elements forming them (Plates 1.26 and 1.27). These multivalent associations most probably resulted from interchanges between the 2 parental chromosome sets at ~~metaphase~~ ^{metaphase} in the hybrid males. As expected in a hybrid between 2 apparently unrelated species, univalent formations were often of a very high order (Plates 1.28 and 1.29), even though the parental types had the same diploid number and it is clear that synapsis failed to occur in many bivalents which might have been considered to be morphological homologous.

Discussion

Karyotypes

The basic chromosome complement or karyotype of a species has proved useful to taxonomists in defining evolutionary relationships

between species within groups naturally defined by morphological and ecological characteristics. Because chromosomes are subject to breakage which may result in the production of inversions and translocations, the karyotype is subject to variation, so that differences between related species which have come about by the accumulation of small changes over long periods may be used to study the evolution of a group. Karyotype evolution may also take the form of changes in the number of sets or the number of sex chromosomes. It is evident from previous studies of the Triatominae that changes have taken place in the basic number of chromosomes and in the sex chromosome complement.

The present study of 13 species of bug, all from South or Central America has revealed a variation in diploid numbers from 21 to 24; but only one species showed reciprocal translocation, all species having the same sex complement. P. pallidus was the only species of the genus of triatomine bug and he concluded that the primitive number was 20+XY for the sub-family. Darlington (1937) suggested that changes in the basic number of chromosomes could occur by means of reciprocal translocations of unequal chromosome sections, producing an increase or decrease in the number. Patterson and Stone (1952) in a study of changes in the group had come about mainly by a reduction in number of chromosomes by centric fusion, inversions and translocations, thus confirming Darlington's (1937) theory. Such previous studies of the Triatominae have been limited to the genus P. pallidus of the Diptera, which provided large numbers of material. In the absence of such material, and with further knowledge of the biology of the group, it remains difficult to say what constitutes the archetypal karyotype of the Triatominae, but since the karyotype of P. pallidus is unique,

Ueshima (1966) is probably correct in assuming that 20 is the basic autosomal complement of the group.

Much greater variation was found in the numbers of sex-chromosomes in the Triatominae studied. It is generally assumed that the XY system represents the primitive condition for sex-chromosomes (John and Lewis, 1965) and the present study has shown 7 species of triatomine with this simple system (T. brevipennis, T. laticollis, T. pallidus, T. pseudo-maculata, T. surdida, T. prolixus and R. neglectus). Six species were found to have multiple sex-chromosome systems, 5 of which were of the type X_1X_2 (T. lenti, T. laticollis, T. pallidus, T. protractus and T. neglectus) and one, T. vittiger, which conformed to $X_1X_2X_3Y$. These X-chromosomes behaved in a regular fashion at meiosis, dividing equationally at the first division and segregating from the Y in a group at the second division. Mann (1958, 1967) has shown that compound X-chromosome systems are typical of the Heteroptera and it is generally thought that the extra X-chromosomes have arisen by fragmentation of an archetypal single X and not by alteration of the autosomes. This hypothesis is strengthened by the demonstration that chromosomal fragments induced by irradiation in species with diffuse systems can give rise to discrete systems, a demonstration which has been confirmed with irradiated T. prolixus. In the present work (Part Four), further suggesting that the multiple XnY system of the Triatominae may have arisen by fragmentation of the archetypal X-chromosome. The demonstration of a constant number of 20 reinforced the view that X-fragmentation is responsible for the XnY system, otherwise a reduction in the number of autosomes would have been observed in the multiple X-chromosome systems.

These species were available for study in the present work, the karyotypes of which had not previously been described- T. leiti, T. tibio-nigra and T. psittaculata. T. leiti had a 20, X_1X_2Y complement while the two T. psittaculata sub-species were 20, X_1X_2Y . Since T. psittaculata and the T. tibio-nigra are all members of the same species, the conclusion of T. psittaculata by Pinto (1966) is severely qualified on karyotype grounds.

Ueshima (1966) concluded from his study of male meiosis in Triatominae that South American species of the genus were of two types, one with an X_1X_2Y complement and the other with an $X_1X_2X_3$ complement, but the present study has demonstrated three South American species (T. leiti, T. tibio-nigra and T. psittaculata) all of which had X_1X_2Y systems. (Since $X_1X_2X_3$ systems are not confined to species in North America as Ueshima (loc. cit.) has suggested.

Female Meiosis

Female meiosis was studied only in a single species, T. psittaculata. The two meiotic divisions, the reduction division with recombination and the second division, were found to occur within about 10 hours after oviposition. It is concluded that the first maturation division takes place either at or just before oviposition, the second division occurring later, between the anterior limit of the common oviduct and sperm enter the micropyle of the egg as they pass into the common oviduct. The first division may occur prior to the formation of the common oviduct, but as most larvae have entry of sperm into the oviduct only the second meiotic division (Wigglesworth, 1965).

Pratt and Davey (1977a) state that work done in their laboratory by Case (1970) showed that the first reduction division of the oocyte nucleus took place in larval T. psittaculata, the age of the larvae not

being specified, but it is clear from the present work that Case (1970) was mistaken and that meiosis, as expected, occurs in adult females only.

Staining of tritamine chromosomes

The present work has demonstrated that G-banding techniques, as developed in recent years for mammalian mitotic chromosomes may be valuably applicable to furthering knowledge about insect cytogenetics. The results obtained with *H. proflus* and *T. tritator* demonstrate that good G-banding can be produced by tritamine chromosomes. A crucial factor in their induction is apparently the initial step of forming a cell suspension. Human cytogeneticists are experienced in preparing cell suspensions in work on blood cultures but insect cytogenetic studies have been restricted to squash techniques which are adequate for the production of G-bands but are not adequate for G-banding which requires separation of cells on the slide and complete spreading on the chromosome. This was achieved in the present study by using the enzyme trypsin with versene to disperse the cells of tritamine embryonic tissue, the resulting cell suspension being fixed, air-dried and banded using the A.S.G. technique of Sumner et al. (1971). The results of this study, together with a previous one, demonstrate the value of mitotic chromosomes within a species, and between species as briefly published by Hardlin (1974). In the present context this is of particular importance to the further study of tritamine bugs, since these insects are particularly well suited to chromosome studies, since chromosomes have been of limited use as taxonomic tools. The polytene chromosomes of the Diptera have, of course, provided numerous genetic markers for that group and fostered sophisticated genetic experiments in relation to the control of Diptera in various situations. It is hoped

that the G-banding technique for trisomic chromosomes described here will provide chromosomal markers for further studies of genetical aspects of problems in the systematics, biology and control of trisomic bugs, besides its interest in insects from South America, possibly for other Hemipteran insects, and perhaps genetical investigations in insects generally.

Hybrid meiosis

Meiosis in male F_1 hybrids from the cross T. maculipes X T. sordidus was characterized by multiple associations and univalent formation, the multiples commonly taking the form of chains of varying size but rings were also found. The production of chains and the multiples suggests that interchanges were produced in the hybrids between some pairs of chromosomes in prophase. The chains produced are similar in morphology to those demonstrated (see later, Part Four) in K. psyllus as a result of interchange formation following irradiation damage. It is difficult to see how such interchanges could persist in holokinetic bugs with low chiasma frequency and linear orientation of chromosomes. The 'invergent' orientation of chain multiples which can result in genetically balanced gametes was not found in these hybrids and the maintenance of ring multiples involving 4 or more chromosomes would have required regular pairing and crossing over which would be difficult given the low chiasma frequency of these bugs. It is not surprising, given the high incidence of univalent and multiple formation, that these F_1 males were found to be infertile and that no F_2 generation could be bred (Petrovagic-Sumelick and Correin, 1972).

The only other cytogenetic study of bug hybrids is by Vaughan (1966) who examined the cross T. sordidus X T. maculipes and found

that the P_2 males typically formed univalents at meiosis. But he did not observe any multiple associations. Further studies of hybrids of such species as could be induced to copulate successfully would probably yield information useful to taxonomists on the basis of the pairing behavior of chromosomes of different species.

P A R T T W O

THE REGULATION OF SPERMATOGENESIS IN MALE

BOONVILLE PROLOGUE

INTRODUCTION

Although it is now well established that hormones play a major role in the regulation of insect growth, the factors responsible for the initiation and control of spermatogenesis in the developing insect remain the subject of debate. Wigglesworth (1940), from his intensive work with *B. proxima*, concluded that there were two major hormonal regulators of growth and differentiation; firstly a moulting hormone (ecdysone) and secondly an inhibitory hormone (allatoin hormone). Wigglesworth (loc. cit.) showed that when female *B. proxima* were given a blood-meal, the resultant abdominal stretching provided a nervous stimulus for neurosecretory cells in the brain to discharge their secretion along their axons to the corpus cardiacum where it was absorbed into the blood as the brain hormone. This brain hormone activated the thoracic gland which in turn secreted the moulting hormone which led directly to the renewal of growth in many organs and tissues of the body, particularly epidermal/cuticular changes associated with the impending ecdysis. The inhibitory hormone secreted by the corpus allatum became operative a few days after the growth and moulting process had already been initiated by the moulting hormone, and for the successive larval stages, prevented the development of imaginal characters. When the final fifth instar *B. proxima* larvae moulted, they undergo a striking metamorphosis and turn into adults with wings and a highly refractive pattern in their cuticle. Wigglesworth (1936) had shown that the moulting hormone, secreted by the thoracic gland, could be modified by implanting the corpus allatum of earlier stage larvae of *B. proxima* into fifth instar larvae; instead of moulting to become adults, the fifth instar larvae moulted to become sixth instar larvae known as supernumerary larvae. Later, Wigglesworth (1940)

suggested that the capacity to form the adult morphology was latent in the larva but was suppressed by the larval genome, but became 'switched on' in the absence of juvenile hormones so that the normal adult form resulted at the final moult.

In holometabolous insects, one instar, the pupa is normally formed between the larva and the adult and metamorphosis starts with pupal formation. The moulting hormone of Holometabola was shown to be similar to the substance found in Hemimetabola, the action being non-specific, including moulting in Hemimetabola and puparium formation in Holometabola (Boggs, 1941). Ecdysone was found to be generally responsible for initiating changes throughout the insect body in those tissues which were limited in growth and differentiation (Boggs, 1944). The juvenile hormone in Holometabola was shown to be produced in small amounts in the last larval stage, which led to the formation of the pupa, while in the pupa no juvenile hormone was formed, allowing the adult to develop.

The life cycle of many insects is interrupted periodically by a phase of arrested development during which metabolism proceeds at a reduced rate, and this arrest of growth is called diapause which is essentially an adaptation to survive adverse conditions. Diapause may be divided into two types, firstly, true diapause which occurs independently of changes in the environment and secondly, periods of quiescence produced by adverse environmental conditions such as cold or starvation. Many factors may be responsible for the induction of true diapause but the most important single factor is said to be day length (de Wilde, 1955). Temperature has also been implicated in the arrest of development of diapausing insects (Boggs, 1944). The larva of B. pallens, when hatched at small size, moulted, but failed to grow because the moulting hormone was not

secreted indicating that growth or moulting could take place in diapausing bugs only if the requisite hormone was given. In a series of experiments with diapausing Lepidoptera, Williams (1942, 1946, 1947 and 1948) showed that the stimulus to diapause in overwintering pupae was the absence of growth promoting factors, the production of which was directed by the brain. Williams (loc. cit.) restored growth to diapausing pupae of the giant silkworm (*Platygonia corrupta*) by chilling, which stimulated the brain to secrete a hormone which in turn activated the prothoracic gland which caused renewed growth by secreting ecdysone.

Spermatogenesis may be considered as a part of the process of insect growth and differentiation and like all other growth processes may be arrested periodically. Using the silkworm moth Williams (1947) showed that pupal diapause could result in the inhibition of spermatogenesis and, since virtually all aspects of insect metamorphosis were known to involve hormones, Schmidt and Williams (1953) investigated the hormonal control of spermatogenesis using in vitro culture techniques. In the Holometabola all the primary spermatocytes are formed by the end of the larval stage but meiosis and spermiogenesis occur in the pupae. Schmidt and Williams (loc. cit.) found that germinal cysts removed from the testes of diapausing larvae would not develop in their culture medium but, when they added blood from metamorphosing silkworms, meiosis began within 24 hours followed by spermatogenesis. Analysis of silkworm larvae haemolymph at different times during metamorphosis revealed a fluctuation in the content of a non species-specific substance, the "macromolecular factor" (MF), which Schmidt and Williams (loc. cit.) suggested was the hormone necessary for sperm maturation. Butenandt and Karlson (1954) found that injection of ecdysone into diapausing

moth pupae rapidly stimulated spermatogenesis in vivo but when ecdysone was added to in vitro cultured testes it had no effect (Karlson, 1956). This paradox was resolved by Karybysellis and Williams (1971a and b) who cultured intact dissecting pupal testes of the silkworm Bombyx mori and showed that both ecdysone and the "macromolecular factor" were essential for in vitro spermatogenesis to proceed. However, removal of the germinal cysts from the testis walls prior to culture showed that spermatogenesis would proceed in the absence of ecdysone and it was concluded that this hormone served only to alter the permeability of the testis wall to the passage of MF which was directly responsible for the induction of spermatogenesis.

The regulation of spermatogenesis in Hemimetabola is different from the Holometabolous process, for in the former some mature sperm may form in the larval stages and spermatogenesis may continue during the adult life. In the silkworm Bombyx mori, which is hemimetabolous, meiotic division starts in the fifth instar larva and proceeds, at a slower rate, in the adult. Economopoulos and Gordon (1971) showed that testes taken from fourth instar larvae of the milkweed bug and transplanted into mature male or female adults underwent extensive spermatogenesis and concluded that moulting was not essential for testis differentiation. Since it was known that a high titre of ecdysone was normally found in fifth instar larvae (Piper and Hsiao, 1957), Economopoulos and Gordon (1971) suggested that spermatocyte differentiation in the fifth instar was stimulated by some change in the blood composition caused by the "adultoid" chemical changes which followed the moult to the fifth instar, and not caused by the direct action of ecdysone on the spermatocytes.

This hypothesis was very similar to that proposed by Kambysellis and Williams (1971) for the regulation of spermatogenesis in holometabolous insects.

The juvenile hormone has also been implicated as a regulatory factor in spermatogenesis. Senhal (1968) implanted corpora allata into pupae of the holometabolous lepidopteran Galleria mellonella and found that testis development was inhibited. Takeuchi (1969) found that spermiogenesis did not occur in testes transplanted from third instar larvae of the silkworm Bombyx mori to pupae with intact corpora allata but did occur in about half of the pupae which had their corpora allata removed, and he concluded that the juvenile hormone had an inhibitory influence on spermatogenesis in the silkworm. In the hemimetabolous cockroach Blattella germanica the testis consists of spermatogonia until the fourth instar when differentiation into primary spermatocytes occurs followed by meiosis and later spermiogenesis which continues in the fifth instar and is completed when metamorphosis to the adult occurs (Amerson and Hays, 1967). Walters and Linsen (1967) found that removal of corpora allata from fifth instar larvae of the cockroach Leucophaea maderae stimulated spermatogenesis in the 'adultoid' insect produced at the next moult and suggested that juvenile hormone normally inhibited sperm differentiation. Economopoulos and Girden (1971) treated fourth and fifth instar Onopeltus fasciatus with synthetic juvenile hormone but could find no changes in the process of spermatogenesis in this blood sucking, contrary to other workers, claimed that juvenile hormone did not play an inhibitory role in spermatogenesis of hemimetabolous insects.

The hemimetabolous haematophagous bug Rhodnius prolixus has five larval stages, in each of which it takes a single large bloodmeal and

then proceeds to moult and grow (Martens, 1971). Wigglesworth (1934) showed that starved R. prolixus larvae would enter a state of diapause which was broken by feeding which led to the release of ecdysone and the renewal of growth. Preliminary investigations for the present work had shown that testis differentiation in R. prolixus was similar to that of the milkweed bug (Economopoulos and Gordon, 1971) in that meiotic divisions started in the fifth instar larva. However, R. prolixus, being haematophagous, has a completely different feeding pattern from the plant-sucking bugs and Schreiber et al. (1968) have suggested that meiosis in tristomatine bugs was initiated by the blood-meal, but did not present any experimental evidence to ~~confirm~~ ^{support} this hypothesis. Recently Dunser and Davey (1974) have investigated the role played by the juvenile hormone in controlling spermatogenesis in R. prolixus and found that they could produce a dose-response related inhibition of the number of spermatid cysts produced by the bug after topical application of juvenile hormone analogue to fifth instar larval males.

The present work was designed to investigate the influence of diapause and blood-meal on spermatogenesis in R. prolixus and in particular to study their influence on intra-meiotic events.

MATERIALS AND METHODS

Materials

All experiments were carried out on R. prolixus from the L.S.H. and T.M. colony, details of which have already been given.

Methods

Bugs were kept in incubators at 25°C (except where specified below) and approximately 70% R.H., in 2" x 3 1/2" glass tubes and fed on rabbit blood.

1. Preliminary experiments

The process of spermatogenesis in normal, untreated, bugs was first examined by making, at daily intervals, testis squash preparations from larval R. prolixus selected at random from groups of bugs of the same age. This was done for fed and unfed fourth and fifth instar larvae.

The testes were dissected out in insect saline and fixed in 3:1 absolute ethanol:acetic acid and stored at 4°C. Squash preparations were made by tapping out testes on to clean slides in a drop of lacto-propionic-orcein, and squashing them under a coverslip, which was then ringed with rubber solution prior to microscopic examination.

2. Timing of meiosis by autoradiography

Groups of male fifth instar larvae and adults, both fed and starved, were injected with an aqueous solution of thymidine- $5-^3\text{H}$ (specific activity 5000 $\mu\text{Ci}/\text{mm}$, Radiochemical Centre, Amersham, England) using a finely drawn glass pipette attached to a micrometer syringe calibrated to deliver known volumes to the nearest microlitre. Each bug was given approximately 1 μCi ^3H -thymidine in 4 μl distilled water injected, via the thin cuticle at the base of the metathoracic legs, into the haemolymph, the puncture being sealed with adhesive wax. For each group of experimental bugs injected with ^3H -thymidine, a control group was injected with an equal volume of water.

Insects were randomly chosen from each group and sacrificed at daily or 12 hourly intervals, testes were dissected out under saline, fixed in 3:1 absolute ethanol:acetic acid and stored at 4°C. Following hydrolysis in 5N HCl for 1 hour at room temperature, testes were stained by the Feulgen reaction and squash preparations of whole testes made in a drop of 45% acetic acid on acid-cleaned slides. Preparatory to

applying the photographic emulsion, coverslips were removed with a razor blade after the preparations had been frozen in liquid nitrogen, and the slides then plunged into absolute ethanol, transferred via 85%, 70%, 55%, 40%, 15% and 10% ethanol to distilled water and allowed to air dry overnight in a dust-free atmosphere.

Autoradiographs were prepared by dipping the slides into a Coplin jar containing 1:1 Nuclear Research Emulsion (1950 Ltd., Essex, England) diluted 1:1 with distilled water, the emulsion being allowed to dry, and the slides then being placed in light-tight boxes for exposure for 7 weeks at -20°C . The autoradiographs were developed in Kodak D19 developer (Kodak, London, England) for 4 minutes and fixed in Kodak Amfix for two minutes. After rinsing and drying, autoradiographs were permanently mounted in Supersal.

3. RNA synthesis during molting

A group of fifth instar male P. prolinus which had moulted from the fourth instar 21 days previously was divided into two groups, one of which was given a blood meal and the other left unfed. Eight days after this feed testes were removed from the larvae by vivisection under insect saline and placed in solid watchglasses containing pre-warmed sterile insect saline with uridine-5,8- ^{32}P (specific activity 53,000 $\mu\text{Ci}/\text{mM}$, Radiochemical Centre, Amersham, England) at a concentration of 100 μCi per 0.1 ml saline (Henderson, 1964). Testes from bugs, in groups of five, were incubated at 27.5°C for $\frac{1}{2}$ hour and 1 hour while control groups were incubated in insect saline. Following incubation, testes were washed in ice-cubes of insect saline and fixed in 3:1 absolute ethanol:acetic acid and stored at 4°C until preparations were made.

Squash preparations for each group were made on clean slides in

45% acetic acid after passing through the alcohol series to distilled water and through the propylene glycol series which were alcoholized. These two slides from each experimental group were incubated for 1 hour at 37°C in a solution of Ellman's reagent (B.D.H., Poole, England) at a concentration of 0.1 mg/ml in propylene glycol (Haskenhale, 1964), rinsed in distilled water and air-dried.

All slides were dipped in G4 nuclear emulsion, exposed for 7 days at -20°C, developed, fixed and rinsed in distilled water for 15 minutes. The slides were then washed in acetate buffer pH 3.6 and stained by the same method with solution of fast blue made up in the same acetate buffer. Following a rinse in two changes of distilled water, the preparations were cleared and mounted in Euparal, ready for examination by light microscopy for evidence of RNA synthetic activity.

RESULTS

1. Preliminary investigations of spermatogenesis in *R. prolixus*

The structure and cytology of the testis of triatomine bugs have been described in detail by Barth (1956a and b) using *T. infestans*, and this is outlined below for *R. prolixus*, supported by observations made for the present work on testes from fourth and fifth instar larvae and adult males of *R. prolixus*, to serve as a foundation for the more elaborate studies which were planned. In the adult, each testis is made up of seven follicles, two of which are much larger than the rest, and each follicle has a germarium at its apex consisting of primordial germ cells. The spermatogonia produced by the germ cells pass down the follicle, divide repeatedly and become covered by a mantle of somatic cells to form a cyst, mature cysts containing up

to 256 spermatogonia (Barth, 1956b). The spermatogonia then become spermatocytes which undergo meiosis.

The successive stages of normal spermatogenesis are illustrated by the photomicrographs (Plates 2.1 - 2.14) of testis squashes of adult male *R. prolixus*. Plate 2.1 shows a gonial cell at mitotic metaphase which eventually leads to the formation of spermatocytes which undergo meiosis starting with prophase leptotene (Plate 2.2). This in turn is followed by zygotene (Plate 2.3) and pachytene (Plate 2.4). Prophase in Heteropteran meiosis is characterized by a 'diffuse stage', following pachytene, during which the autosomes lose their affinity for nuclear stains but the sex-chromosomes (X₁) remain heteropycnotic; this stage is shown in Plate 2.5. The 'diffuse stage' is followed by a normal diplotene (Plate 2.6), diakinesis (Plate 2.7) and the first spermatocyte division (Plate 2.8) at which point the sex-chromosomes are heteropycnotic and the autosomes have become achromatinized. The first meiosis is followed by anaphase and telophase I (Plate 2.9) in which the sex-chromosomes lie at the centre of a ring of autosomes, and the second meiotic division follows immediately with metaphase II (Plate 2.10), anaphase II (Plate 2.11) and telophase II. There then follows the process of spermatogenesis by which the rounded primary spermatids (Plate 2.12) are elongated into elongated secondary spermatids (Plate 2.13) and finally into mature elongated spermatozoa (Plate 2.14).

Using this knowledge of the stages of spermatogenesis, a preliminary comparison was made of the testes of fed and unfed fourth, and the unfed fifth instar male locusts, and of adult adults allowed to moult through from fed fifth larval bugs. Methods for squash preparation and staining with orcein were as given above (Part One).



Plate 2.1. Gonial mitosis from E. prolixus testis at metaphase.
X 2500



Plate 2.2. Leptotene nucleus from E. prolixus testis. X 2500.



Plate 2.1. Cytogene nucleus from S. malinus testis. Note distinct heteropycnotic X and Y. X 200x.



Plate 2.2. Cytogene nucleus from S. malinus testis. X and Y visible heteropycnotic. X 200x.



Plate 2.5. 'Diffuse stage' of meiotic prophase from E. prolixus testis.
Note euchromatic, diffuse autosomes. X 2500.



Plate 2.6. Diplotene spermatocyte from E. prolixus testis. Autosomes
now more deeply stained. X 2500



Slide 2.7. Spermatocyte at diakinesis from *S. latipes* testis.
X 2500.



Slide 2.8. Metaphase I spermatocyte from *S. latipes* testis. Autosomes
were more deeply stained than sex-chromosomes. X 2500.

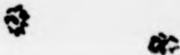


Plate 2.9. Telophase I from H. prolixus testis. Note sex-chromosomes surrounded by a ring of autosomes. X 2500.



Plate 2.10. Metaphase II from H. prolixus testis. Note X and Y in centre of a ring of more deeply staining autosomes. X 2500.



Plate 2.11. Anaphase II from R. prolixus testis. X 2500.



Plate 2.12. Bounded primary spermatids from R. prolixus testis. X 2500.



Plate 2.13. Primary and secondary spermatids from B. prolixus testis.
X 2500.



Plate 2.14. Mature spermatozoa from B. prolixus testis. X 2500.

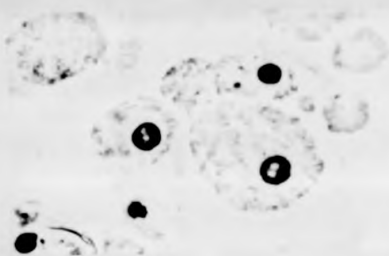


Plate 2.15. 'Aborted' spermatids from R. prolixus male in diapause.
X 2500.

Unfed fourth instar larvae. A large group of fourth instar larvae which had all moulted to this instar on the same day were sampled daily, commencing 14 days after the moult and continuing for 30 days, no blood-meal being given. Two insects were sampled on each day and testis squashes prepared from each one. These testes were all very small on dissection and the squash preparations showed that spermatogenesis had not been initiated in these unfed bugs; not even gonial divisions, as late as 44 days after the previous ecdysis.

Fed fourth instar larvae. A group of fourth instar larvae, all of the same age, were fed on rabbit blood on the same day and sampled at daily intervals for 7 days. At day 3 the first gonial mitoses were detected in squash preparations and these continued up to day 7 with no more advanced stages being seen up to that time. The bugs moulted about the 12th day.

Unfed fifth instar larvae. A group of fifth instar larvae was fed at fourth instar on the same day and those bugs which moulted to the fifth instar, 12 days later, were kept unfed until, at 28 days after the moult (experimental day 0), the testes were dissected and squash preparations made from 5 of them. This was repeated with 5 more of the bugs on every other day up to experimental day 40. On microscopic examination, the follicles of these diapausing larvae were found to be populated largely by two cell types, firstly diffuse stage spermatocytes (Plate 2.5) and secondly what appeared to be 'aborted' primary spermatids. Very few cells in the 1st or 2nd meiotic divisions were seen in these preparations compared with normal testes from fed adults. Secondary spermatids and mature spermatozoa were completely absent from all these larval testes. The 'aborted' primary spermatids were of particular interest and their numbers appeared to increase

proportionately as the period of diapause lengthened. Plate 2.15 shows 'aborted' spermatids photographed in a squash from the testis of a diapausing fifth instar larva. These spermatids were more densely stained than normal (see Plate 2.12) and appeared to undergo a process of degeneration during which their volume increased greatly (Plate 2.11). These results suggested that primary oocytes in diapausing larvae were liable to become hypertrophied, and some either degenerating passively or being actively disposed of by the insects.

2nd Fifth Instar Larvae and Pupae of *M. prolixus*. A large group of male larvae was fed on fourth instar medium on the same day and allowed to moult into fifth instar larvae. These were next fed 28 days after the moult (experimental day 1). After this meal 2 bugs were selected from the group and placed, respectively, under 1st and 2nd testes, and 2 bugs were dealt with in this way daily for a further 24 days, on into the adult stage into which they moulted in about 18 days. Examination of these squashes revealed great differences between these testes and those of the unfed diapausing fifth larval bugs. In the fed bugs 1st and 2nd meiotic divisions were abundant by 3 days after the blood-meal. This wave of meiosis was followed on day 10 by the first signs of spermatogenesis and on day 12 the first mature spermatids were seen.

These bugs moulted to the adult stage about day 18 after the blood-meal and examination of the adult testes continued for 6 more days during which time they were still actively undergoing meiotic divisions and spermatogenesis. Aborted oocytes which were found in unfed fifth instar larvae were not seen in fed fifth instar larvae after the first 3 days following the blood-meal.

Having completed these preliminary investigations of spermatogenesis in *M. prolixus* the following conclusions were drawn:-

1. Testis differentiation commences in fourth instar males following blood-meal.
2. Spermatocyte production does not start until the 1st moult into fifth instar.
3. Spermatogenesis will not proceed in diapausing 5th instar larvae and the primary spermatids produced develop no further and 'abort' during diapause.
4. Following a blood-meal, fifth instar larval spermatocytes rapidly undergo meiosis and the process of spermatogenesis is inaugurated. Some 'aborted' primary spermatids are evident only for about 3 days after the blood-meal and these are, presumably, those which form during the pre-feed period, in this case 28 days.

2. Timing of meiotic

Because of the apparent importance of the blood-meal to the process of spermatogenesis, autoradiographic experiments were carried out to time the process in both fed and unfed bugs to determine if the blood-meal affected the rate of intra-meiotic events. The regulation of the meiotic process, whilst of scientific interest, must also be considered of importance in relation to the control of this vector species. The inhibition of spermatogenesis may be a means of producing sterile males for population control purposes.

The principle involved in these experiments is well established.

It involves simply recording the most advanced cell stage which is labelled at different time intervals after a 'pulse' label of ^3H -thymidine is given, the first labelled cells to reach a particular

meiotic stage being those which incorporated the thymidine at the end of the preceding DNA replication period or S-phase (Honema, 1962; Mackenthauer, 1964; Taylor, 1965; Callan and Taylor, 1968; Moens, 1970; Coggins and Gall, 1972). Selman and Kafatos (1974) have recently shown that nearly all ^3H -thymidine is incorporated into DNA when injected into insects and after a single injection is rapidly cleared from the haemolymph, resulting in a maximum 'pulse' of 4 hours. The sampling interval between injection and examining insects on the first occasion and subsequently in the present experiment, was a minimum of 12 hours which in view of the work of Selman and Kafatos (loc. cit.) must be given an error margin of 4 hours. In scoring labelled slides, comparison was made throughout with control autoradiographs from bugs injected with water.

Fed fifth instar larvae

(a) At 25°C. Bugs used in this experiment were fed simultaneously as fourth instar larvae and again as fifth instar larvae 28 days after the fourth instar feed. The bugs were then injected 7 days after the blood-meal (experimental day 0). The rate of progress of cells through the process of meiosis for fed fifth instar larvae kept at 25°C is shown in Fig. 2.1 in which each line represents the stage(s) found labelled within a single insect. It may be seen from this figure that meiosis was completed in 12 days at this temperature, the longest stage being the 'diffuse stage' which lasted about 4 days.

(b) At 27.5°C. Bugs for this experiment were of the same age and received the same treatment as fed fifth instar larvae kept at 25°C. The results of this experiment are given in Fig. 2.2 which show that at this higher temperature meiosis was completed in 10 days.

In a view from a comparison of Figs. 2.1 and 2.2 about meiosis

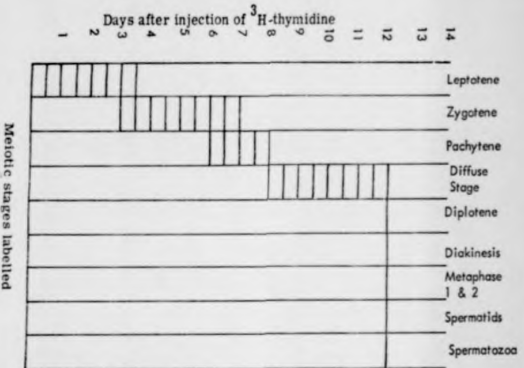
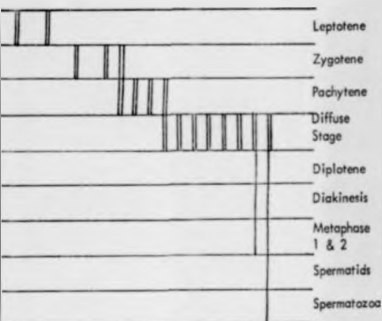


Figure 2.

Timing of meiosis in *Rhodnius prolixus* fed fifth instar larvae at 25°C. Each line represents the stages labelled within a single insect.

Days after injection of ^3H -thymidine

2 3 4 5 6 7 8 9 10 11 12



environmental temperature shortens the time spermatocytes take to pass from early prophase to mature spermatozoa. As at 25°C, so for 27.5°C, spermatocytes on leaving the 'diffuse stage' passed through diplotene, diakinesis, both meiotic divisions and spermiogenesis in less than 24 hours. Fig. 2.1 shows that the first labelled metaphase II cells were seen on day 10 but the first labelled spermatozoa were not seen until day 10½ which suggests that spermiogenesis can occupy about 12 hours at this temperature.

Unfed fifth instar larvae

The bugs used in this experiment were all fed as fourth instar larvae on the same day, allowed to moult to fifth instar larvae but these were not fed. Experimental day 0, when the ³H-thymidine was injected, was 35 days after the fourth instar blood meal was given, and approximately 21 days after the moult to the fifth instar. This long period of starvation ensured a state of diapause (Wigglesworth, 1934).

The results of this experiment are given in Fig. 2.3 which shows that, in diapausing fifth instar larvae, meiosis takes approximately twice as long as in fed fifth instar larvae at the same temperature. Meiosis took as long as 20½ days in the unfed fifth instar larvae compared with only 12 days in fed larvae. Healthy spermatids and mature spermatozoa were not found in the testes of unfed bugs and the results show that the first labelled 'aborted' spermatids appeared on day 21. Whilst the prophase stages of leptotene, zygotene and pachytene occupied approximately the same time periods in fed and diapausing larvae, the 'diffuse stage' lasted from day 11 to day 20 in unfed bugs (a period of 9 days) while in fed bugs it occupied only 4 days.

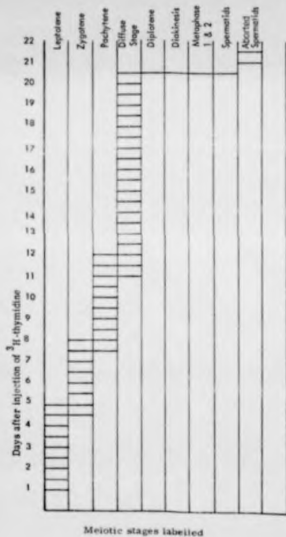


Figure 2.3. Timing of meiosis in Rhodnius prolixus using fifth instar larvae at 25°C . Each line represents the stages labelled within a single insect.

Fed adults

Adult males, which, as fifth instar larvae had been fed on the same day, were fed again as adults 47 days later, and then injected with ^3H -thymidine 7 days after this adult blood-meal (experimental day 0). The results (Fig. 2.4) show that the successive stages occupied approximately the same time periods in fed adults as they did in fed fifth instar larvae (Fig. 2.1). The process of meiosis was completed in $12\frac{1}{2}$ days in fed adults. Some of the labelled stages in this experiment are shown in photomicrographs on Plates 2.16 - 2.21. Plate 2.16 shows a labelled zygotene cell from a fed adult and the other labelled stages shown are pachytene (Plate 2.17), 'diffuse stage' (Plate 2.18), metaphase I (Plate 2.19), anaphase II (Plate 2.20) and Plate 2.21 shows labelled spermatozoa.

Unfed adults

The males used in this experiment were fed as fifth instar larvae on the same day as in the above experiment with fed adults, but they were not given a blood-meal after moulting to adults. They were injected with ^3H -thymidine 47 days after feeding as fifth instar larvae (experimental day 0) and approximately 35 days after they had moulted to adults.

The results of this experiment are given in Fig. 2.5 which shows that meiosis occupied 16-17 days in these diapausing adult males as opposed to $12\frac{1}{2}$ days for fed adult males. Some labelled spermatozoa as well as labelled 'aborted' spermatids were found in the unfed male adults, suggesting that spermatogenesis is not completely inhibited in diapausing adults. Nevertheless, the fact that no labelled metaphase divisions or spermatids were seen in these preparations (Fig. 2.5) suggests that few cells were passing through division or

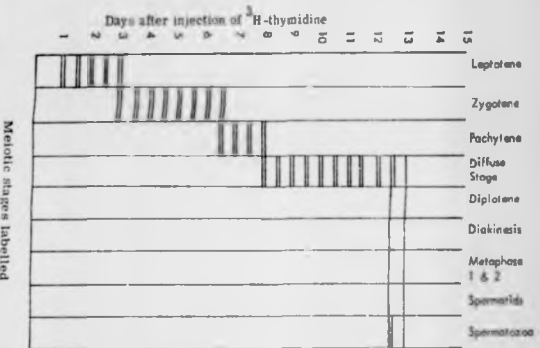


Figure 2.4.

Timing of meiosis in Anopheles prolixus fed adults at 25°C.
Each line represents the stages labelled within a single insect.

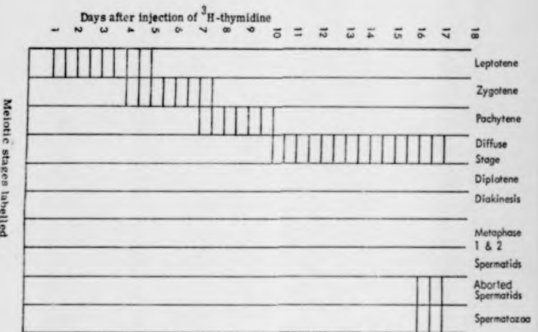


Figure 3.2.

Timing of meiosis in Rhodnius prolixus unfed adults at 25°C.
Each line represents the meiosis labelled within a single
insect.



Figure 2.16. Automerism of 1980-1981 H. prolixus testes showing
with ^3H -thymidine showing zygotene nucleus. X 2500.

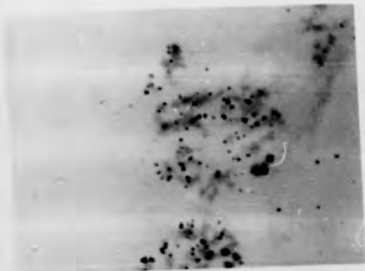


Plate 2.17. Autoradiograph from adult H. prolixus testis labelled
with ^3H -thymidine showing labelled pachytene nucleus.
X 2500.

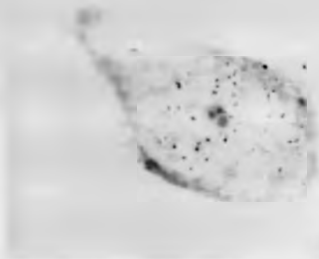


Figure 2.0. R. Autoradiograph from adult R. prolixus testis labelled with ^3H -thymidine showing labelled 'diffuse stage' nuclei. X 2500.



Figure 2.1. R. Autoradiograph from adult R. prolixus testis labelled with ^3H -thymidine showing labelled metaphase I nucleus. X 2500.

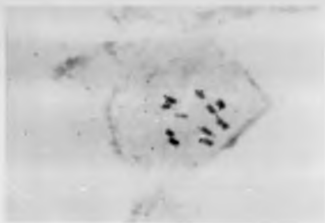


Plate 2.20. Autoradiograph from adult *S. prolixus* testis labelled with ^3H -thymidine showing labelled metaphase II nucleus. X 2500.



Plate 2.21. Autoradiograph from adult *S. prolixus* testis labelled with ^3H -thymidine showing labelled spermatozoa. X 2500.

spermiogenesis and that inhibition of these processes was severe in these diapausing adults.

As in the unfed fifth instar larvae, the 'diffuse stage' was greatly extended in these starved adult males, taking about 7 days compared to 3 days for starved adult females.

In summary, these autoradiographic studies show that the rate of reduction in *P. pallidus*, judged by the first appearance of successive labelled stages, varies according to the nutritional status of the bug. Thus, at the same temperature, the fifth instar larval spermatocytes took about 12 days to pass through meiosis, while in unfed larvae the process occupied about 21 days, this difference being accounted for mainly by variation in the amount of time cells spent in the 'diffuse stage'. The rate of meiosis appeared to be very similar in fed adults and fed fifth instar larvae, both groups taking about 12 days. However, cells of diapausing larvae took 5 days longer to complete meiosis than those of diapausing adult cells, which may be related to the degree of starvation of the two groups.

3. RNA synthesis during spermatogenesis

Because of the difference found between fed and unfed bugs in the time taken to pass through the different spermatogenic stages, it was decided to compare the process of RNA synthesis in fed and unfed males to see if differences could be detected in cell synthesis related to the nutritional status of the bug.

The incorporation of ³H-uridine into the nuclei of spermatocytes of fifth instar larvae was examined by grain counts made on the nuclei of unextracted squash preparations made from the testes of fed and unfed fifth instar larvae incubated for $\frac{1}{2}$ and 1 hour in the

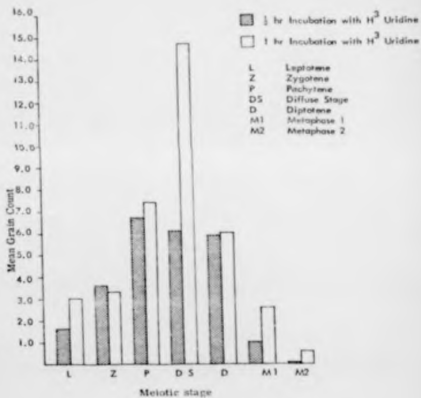


Figure 2.6. RNA synthesis in *Rhodnius prolixus* fed fifth instar larval testis.

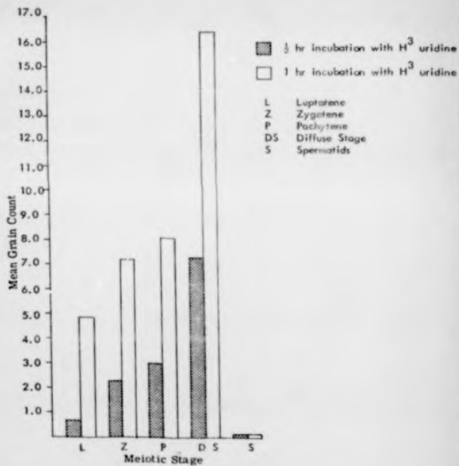


Figure 2.7. RNA synthesis in *Rhodnius prolixus* unfed fifth instar larval testis.

labelled precursor. The slides extracted with RNA-ase showed no evidence of any labelling and did not differ from untreated control slides, which confirmed that the labelled slides were indicating synthesis of RNA.

Grain counts were carried out over 10 cells of each meiotic stage from each preparation, both control and labelled, and the mean cell count was calculated by taking the mean count for 10 labelled cells of any one stage and subtracting the mean score of 10 cells of the same stage in the control slides. In practice, the counts from the control slides were very low, often scoring zero.

The results of this experiment are shown in Fig. 2.6 for fed fifth instar larvae and Fig. 2.7 for unfed fifth instar larvae.

These figures show that the labelling patterns were very similar for both the $\frac{1}{2}$ and the 1 hour incubation periods. Throughout meiotic prophase spermatocyte nuclei were actively synthesising RNA in both fed and unfed larvae. In fed and in diseasing larvae the 'diffuse stage' nuclei were found to be the most heavily labelled and by inference the most actively synthetic. High levels of nuclear labelling continued up to diplotene in the spermatocytes of fed larvae and the level of labelling fell off rapidly in cells engaged in the meiotic divisions. There was no evidence of any RNA synthesis during spermatogenesis in these fed bugs.

In the testes of ⁴⁵Su larvae (Fig. 2.8) cells in the diplotene, diakinesis or metaphase stages were not found and, as in the studies of ³H-methylidine incorporation, there appeared to be no spermatogenesis taking place in these testes and the 'aborted' spermatids showed no evidence of synthesising RNA.

Plate 2.22 shows a 'diffuse stage' spermatocyte labelled with

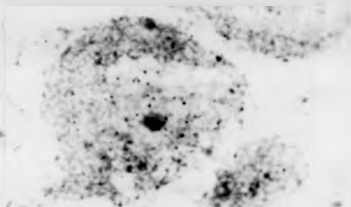


Plate 4.22. Autoradiograph from dissection of fifth instar B. morio testis labelled with ^3H -uridine showing labelled 'abortive stage' nucleus. X 2500.

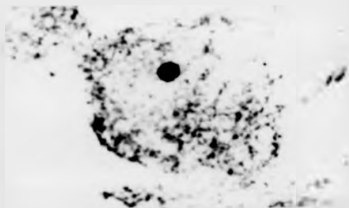


Plate 4.23. Autoradiograph from dissection of fifth instar B. morio testis labelled with ^3H -uridine showing unlabelled 'aborted' spermatid. X 2500.

³H-uridine from the testes of a diapausing larva after 1 hour incubation in the precursor and the high level of label indicating synthetic activity is readily observable. Figure 7.13 shows an 'aborted' spermatid from the same bug which indicates that there was no RNA synthesis taking place in these nuclei. These results suggest that the elimination of 'aborted' spermatids may be a passive process in the diapausing bug involving little or no nuclear transcription of RNA.

DISCUSSION:

A preliminary examination of squash preparations taken from testes of fourth and fifth instar male larvae killed sequentially revealed that mature flagellated spermatozoa were first produced in fifth instar larvae after a blood-meal. In unfed fifth instar larvae, spermatocytes passed through meiosis and rounded primary spermatids were produced, but spermiogenesis, the process by which spermatids are converted into flagellated spermatozoa (Wilson, 1975), did not take place and instead primary spermatids became densely stained and apparently 'aborted'. Spermiogenesis was rapidly initiated in diapausing fifth instar larvae given a blood-meal so that prior to the moult to the adult form, R. prolixus males were found to have a supply of mature spermatozoa. These preliminary investigations were in close agreement with the results of Dummer and Davey (1974) who studied sections of R. prolixus testes and who further suggested that 'aborted' spermatids which they found in diapausing bugs were removed by autolysis.

The critical factors affecting production of mature spermatozoa in R. prolixus appeared to be age and nutritional status; age being

critical since testis differentiation would not take place in fourth instar larvae, and the blood-meal being critical since spermatogenesis would proceed only as far as the production of primary spermatids in diapausing fifth instar larvae. This critical effect of age is similar to that demonstrated by Economopoulos and Gordon (1971) in the milkweed bug testis which they attributed to chemical change in the insect associated with age. The critical sensitivity of testis development to bug age suggests the influence of a genetic mechanism in B. pumilus switching on the process of differentiation in a manner analogous to that proposed by Wigglesworth (1968) for other adult structures in B. pumilus. Conner and Wang (1974) have suggested that the juvenile hormone is responsible for the inhibition of spermatogenesis in B. pumilus and my suggest for the age dependency of the process, since secretion of juvenile hormone is prevented in fifth instar B. pumilus by sexual control from the male (Wigglesworth, 1968). However, the present work shows that fast-diapausing fifth instar larvae will not produce mature spermatozoa and therefore some signal or product associated with the blood-meal is necessary for spermatogenesis to be completed. The present experiments do not reveal the nature of such a signal; it may be that the situation in B. pumilus parallels that in Molometabolis in which a blood-borne macromolecular factor (MF) is apparently required to break the diapause inhibition of spermatogenesis in silkworm pupae (Kambyse'lis and Williams, 1971a and b).

Although the precise nature of the blood-meal stimulus to spermatogenesis was not determined, the effect of the blood-meal on the rate of spermatogenesis has been investigated by meiotic timing techniques using ^3H -thymidine label. Spermatocytes of fed fifth instar larvae kept at 25°C passed through meiosis in approximately 12 days while at 15°C the process was prolonged to 20 days. The results suggest

between temperature and rate of meiosis is well documented for other insects (Craig-Cameron and Jones, 1970) and was not of primary interest in the present experiments except to note that all comparative studies were carried out at 25°C. A further experiment showed that meiosis from leptotene to primary spermatid stage took approximately 21 days in diapausing fifth instar larvae, 9 days longer than in fed larvae, a difference which was largely accounted for by the time taken for spermatocytes in the unfed bugs to pass through the 'diffuse stage' of meiosis. Similar results were recorded for fed and diapausing adult males in which the 'diffuse stage' was extended from 4 days in fed bugs to 7 days in unfed bugs. Although the same period of diapause was imposed on both fifth instar and adult bugs (35 days), diapausing larvae took 5 days longer to complete meiosis than did diapausing adults. Fifth instar larvae R. prolixus take a massive bloodmeal of about 250 µg which is more than twice the amount taken by fourth instar larvae (Burton, 1930) so that adult bugs may have derived greater food reserves from the fifth instar than those derived from their fourth instar after a similar period of starvation. This may account for the difference in rate of meiosis between starved fifth instar larvae and starved adult males.

Dummer and Davey (1974) found that application of juvenile hormone analogue to R. prolixus produced a dose-related inhibition of meiosis which they assessed by the number of spermatid cysts present in sections of testis. They found that the size of the 'spermatocyte containing' part of the testis remained constant and concluded that the effect of the juvenile hormone had been on the rate of gonial mitoses and not on the meiotic process itself. From these experiments Dummer and Davey (loc. cit.) proposed a kinetic hypothesis for the

regulation of spermatogenesis in P. prolixus, which assumed an endogenous level of production in the absence of endocrine activity which would be increased by ecdysone following the blood meal, to produce a high rate of cell division and differentiation which could be returned to the endogenous rate by the secretion of the inhibitory juvenile hormone. However, this hypothesis was based on the premise that the duration of meiotic prophase was a constant and that only the rate of gonial input varied. The present experiments have shown that meiotic prophase is not of fixed duration in spermatogenesis and may vary in length considerably depending on the nutritional status of the bug, the flexibility lying in the amount of time spent in the 'diffuse stage'. The 'diffuse stage' of meiosis in the suborder Heteroptera (Order Hemiptera) (Lewis and Gudder, 1958) and is associated in P. prolixus with a great increase in nuclear volume (Schreibler et al., 1961) indicating that chromosomes undergo a process of the condensation they have gone through up to pachytene and lose their affinity for nuclear stains. The ability of spermatocytes in this 'diffuse stage' to respond to changes in the internal environment of the bug is of considerable interest since, in many other animals, ~~spermatocytes~~ ^{spermatocytes} have been demonstrated in nuclear behaviour at the prophase-diplotype stage of meiosis. For example, in grasshoppers meiosis is typified by lateral 'lampbrush' loops which extend from the chromosomes during pachytene and a prolonged diplotene, but prior to first metaphase the lateral loops are withdrawn and compact chromosomes are formed (Hill and Hill, 1961). There is evidence that 'lampbrush' chromosomes are widespread throughout the animal kingdom in meiotic cells of both sexes (Hill, 1961, 1962). For example, Hill (1961) and Hill has demonstrated 'lampbrush' chromosomes in the oocyte nuclei

of grasshoppers, locusts and cockroaches and there is evidence that chromosomes pass through a 'lampbrush' stage during male meiotic prophase in Drosophila (Meyer, 1961; Callan (1967) has suggested that the pattern of chromosome organisation in eukaryotes consists of a series of repeated sub-unit sequences, each made up of a 'master' followed by 'slave' sequences and that the extension of the lateral loops during 'lampbrush' formation results from the progressive matching of 'slaves' against the 'master' copy thus correcting any errors in the 'slaves' which are first formed and once the lateral loops, full of 'slave' sequences have grown to their greatest extent in meiotic cells, these are then removed. This suggests an additional stage follows the process of genetic recombination in meiosis and, following correction, the 'slaves' are then to be destroyed if required. There is evidence that RNA synthesis takes place in the lateral loops of 'lampbrush' chromosomes of crickets (Gall and Callan, 1962) and during 1967 we found that the 'lampbrush' loops of the Y-chromosomes of Drosophila spermatocytes are associated with intense RNA synthesis.

Although 'lampbrush' chromosomes have not been demonstrated in Salmonella but, the 'Hirtle stage' of male meiosis, which is an extended pachytene/diplotene phase, may represent a system of chromosome organisation analogous to Drosophila formation and serving similar functions. The observation of Stomatopus sp. in the testes of R. n. p. l. x. demonstrated in the present work has shown that 'Hirtle stage' spermatocytes are full of RNA synthesising lateral loops extending distally (cf. the pattern of meiotic prophase has been shown to be very different in other insects (Hirtle, 1961; Meyer and Pech, 1961 and Pech 1961). Call suggested these meiotic stages

produced during meiosis may carry information related to the control of subsequent steps of spermatogenesis. This hypothesis was confirmed in part by Haas (1965) who showed that Drosophila males which lacked a single Y-chromosome 'Jambbrush' loop were sterile. This sterility resulted from the arrest of spermiogenesis at different stages depending on which loop of the chromosome was missing, suggesting that loop products were released at first metaphase specifically to control spermiogenesis.

The present experiments have shown that spermiogenesis is inhibited during diapause in both larval and adult R. prolixus and ³H-thymidine labelling has shown that the blood-meal has a remarkable accelerative effect, apparently acting directly on 'diffuse stage' spermatocytes. The results suggest that a blood-borne factor is released following the blood-meal and it may be that a 'macromolecular factor' similar to that demonstrated by Thompson and Williams (1971a and b) is involved. Pratt and Davey (1972a, b and c) have shown that egg production in R. prolixus females may be affected by many factors, including hormones and starvation, which may be seen as an adaptation to the haemophagous way of life, eggs being produced in a cyclical fashion each feed results in a cycle of egg production, the number of eggs depending on the size of the meal ingested. It could be postulated that the production of sperm in the male is similarly governed by the nutritional status of the bug. If this were the case, it could further be postulated that a blood-borne factor released after a blood-meal interacts directly with the uncondensed 'diffuse stage' spermatocyte chromosomes and that the RNA synthetic activity demonstrated in these nuclei was in part associated with control of the subsequent stages of spermatogenesis and, in particular, spermiogenesis, in a way analogous to the 'Jambbrush' control of meiosis demonstrated for Drosophila males (Haas, 1965).

It is known that insect hormones can act directly on chromosomes, for example Clever and Karlson (1963) induced puffs in the giant chromosomes of Chironomus tentans larvae with ecdysone and it has been shown that the hormone acts directly on the nucleus to induce RNA and in turn enzyme synthesis (Karlson, 1961, 1963 and 1967). Considering this information from other organisms, it is reasonable to propose that a hormonal stimulus (possibly a 'macromolecular factor') acts as a switch to turn on r-RNA synthesis in the 'diffuse' chromosomes of the spermatocytes of diapausing K. prolixus, thus providing the biochemical stimulus for spermiogenesis to proceed after the blood-meal. However, since it has been shown in the present work on diapausing unfed adult males that at least a few cells pass from the 'diffuse stage' and through spermiogenesis, it may be that the control substance is quantitative in its effect, depending on the degree of starvation of the bug.

PART THREE

INHERITANCE OF SUSCEPTIBILITY TO TRYPANOSOMA CRUZIINFECTION OF PEROMYSCUS PROLIXUS

INTRODUCTION

The clinical diagnosis of Chagas' disease is difficult in the chronic phase, the persons may harbour trypnosomes all their lives yet may be symptomless and acting as carriers (Marden et al., 1969). Even in acute cases the number of trypnosomes in the blood may be too few to be detected by direct methods and other means must be employed such as immunodiagnosis. The various immunological tests for Chagas' disease have recently been reviewed (Goble, 1970) and the oldest of these, the complement fixation test, first used by Guazirero and Machado (1913) only 4 years after the discovery of T. cruzi, is still the most widely used serological test.

Immunological tests rely on sophisticated techniques and yield 'false positive' diagnoses; the simplest ~~unpleasant~~ expensive way of diagnosing chronic Chagas' disease is by the method of xenodiagnosis which does not produce 'false positives' and of all the diagnostic tests available is the most sensitive (Wander, 1970). Xenodiagnosis was first described by Brumpt (1914) as a method for using the natural vector of a disease to diagnose an infection in a host animal. The procedure is simple, usually 6 fifth instar larvae from a clean laboratory colony are allowed to feed on the patient and the bugs are then examined some time later by either pulling out the gut and examining the contents or by compressing the abdomen causing involuntary excretion and then examining the faeces (Dias, 1934b). Maskitt (1964) suggested an improvement to the technique which involved homogenizing the test bugs in saline, filtering, centrifuging and examining the sediment for trypnosomes. Comparing this method with the squash technique on the same group of patients, Maskitt (loc. cit.) found a significant improvement in the sensitivity of the test. However, the squash technique,

whereby the gut contents are simply pulled out on to a slide, diluted with saline and examined for trypomastotes, is still the most widely used method (Siqueira, 1968). The main objection to xenodiagnosis is its inefficiency in detecting very low levels of parasites; for example, Pizano (1954) using laboratory animals with known chronic infections, found that Triatoma protracta detected 44% of the positive cases in guinea-pigs and 77% in dogs. Moreover, within the groups of bugs tested on each positive animal, only 50% were individually positive. Romero and Romana (1957) in an experiment on 433 patients showing clinical symptoms of Chagas' disease found that only 30% were positive by xenodiagnosis while 96% were positive by complement fixation test. De Freitas (1961) using fifth instar Triatoma protracta obtained only 20% positive results on patients with chronic Chagas' symptoms and Marsden et al. (1969) found that 16 out of 20 Triatoma protracta fed on patients with a patent parasitaemia failed to develop flagellates. Clearly, estimates of the efficiency of xenodiagnosis vary widely and the present work was designed to investigate the extent to which the insect genotype affects this variation and if so, whether the efficiency of the insects as diagnostic tools could be improved.

Many factors may influence the development of a Trypanosoma infection in a bug after the infecting blood-meal at which the vector ingests the parasite. A bug which will support the development of the parasite is said to be 'susceptible' and in the same way the term 'refractory' describes individuals or populations in which the parasite will not develop (Macdonald, 1967). 'Intensity of infection' is used to describe the numbers of parasites which the vector will support. Whether or not a bug is susceptible, and the intensity of this infection, both affect the vector efficiency, which will influence the choice of

bugs for serodiagnosis. Vector efficiency may be affected by both bug and parasite characteristics, environmental factors and interactions between these (see Fig. 3.11). The most important environmental factors which influence the physiology of insects are humidity and temperature (Wigglesworth, 1965). Buxton (1932) showed that adult H. pallidus possess considerable powers of regulating the proportion of water in their bodies and that only prolonged desiccation (42 days at less than 60% R.H. and 23°C) would cause these regulatory processes to break down. Desiccation results eventually in body injury and fall in metabolic rate and it is therefore important to ensure humid conditions in keeping bugs experimentally (Buxton, 1932). Estimates of optimal conditions of humidity vary, Ryckman and Ryckman (1966) recommended 50-60% R.H. at 27°C for Triatomines while Zolodov et al. (1967) suggested 70-80% R.H. at 23°C. Temperature affects many aspects of the bug-trypanosome relationship since, as the external temperature rises, so the metabolic rate of insects rises. The relationship between temperature and the rate of biological processes is described by an S-shaped logistic curve (Davidson, 1942, 1944) which represents the speed of development of many insects over most of the temperature range which will support development. The rate of development of a bug from a first instar larva to the adult is, of course, a function of metabolic rate and in turn temperature, given that appropriate blood-meals are taken between moults. (Triatomine bugs have 5 larval instars 1/2 each of which they take one or more large blood-meals and then moult to the next instar). Buxton (1912a) found metacyclic trypanosomes in the bug Triatoma sanguinolenta at varying time intervals following an infecting blood-meal, depending on the temperature at which the bugs were kept and Phillips (1960b) found that the interval between an infecting meal and evidence

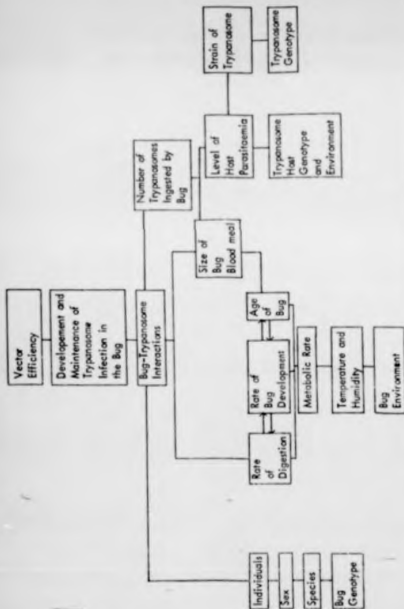


Figure 3.1. Factors affecting vector efficiency

of trypanosomes appearing in the bug faeces decreased progressively as the temperature increased in bugs of all larval stages. Wood (1954) noted that adult T. prolixus caught during the summer in California had more trypanosomes in their faeces than bugs caught in mid-winter and also showed that increasing the temperature increased the number of trypanosomes in the faeces of experimentally infected bugs.

Since temperature affects the rate of development of a bug, the age of a bug may be regarded as a function of temperature and several studies have examined the effect of age on the trypanosome relationship. Buxton (1931) showed that he could infect bugs at all ages experimentally, and in a survey of wild caught T. mexicanus Lorie (1915) noted that the proportion of infected bugs increased with age, which he attributed to the fact that older bugs had taken in more blood and would have had more opportunities for reinfection. By counting the number of trypanosomes in faecal droplets, Wood (1954) found that adult T. prolixus excreted more trypanosomes than larvae over a 6 hour period. Phillips and Bertram (1967) compared the percentages of bugs of each larval instar of T. prolixus which were positive for metacyclic trypanosomes in their faeces after an infecting blood-meal and found that the proportion declined with each larval instar, so that 92% of first instar larvae were infected while only 77% of fifth instar larvae were positive, although adults showed an increase to 88% positivity. In view of these results several factors must be noted; firstly, bugs take increasingly larger blood-meals at successive larval stages ranging from about 6 mg in first instar T. prolixus to about 280 mg in fifth instar larvae and falling to about 170 mg in the adult (Buxton, 1931); secondly, rate of digestion of the blood-meal increases as the bug ages,

the fifth instar larva digesting its meal 20 times faster than the first instar larva (Buxton loc. cit.). Thus, if bugs are fed experimentally on an animal with a known parasitemia, the more blood a bug digests the more parasites it will ingest. Phillips and Bertram (1967) estimated that first instar larvae ingested approximately 40,000 trypanosomes while fifth instar ingested over a million trypanosomes when fed on rats with known parasitemia and yet the percentage remaining infected decreased with age. Wood (1954) also noted that bugs which received fewer parasites with their infecting meal showed as many parasites in their feces as those ingesting large numbers of trypanosomes. Recently Smith et al. (1968) have shown that the rate of infection was higher in later stages of experimentally infected H. prolixus, but their data shows that this trend was not consistent between replicate experiments and that in one experiment second instar larvae showed the highest infection rate and fifth instar larvae the lowest.

From these results it appears that neither the size of the blood-meal nor the number of trypanosomes ingested determines whether or not a bug will maintain an infection, or the intensity of that parasitemia. Phillips and Bertram (1967) suggested that the increased rate of digestion of older bugs was harmful to the development of ingested trypanosomes and Dias (1934a) also related parasite survival with the rate of hemolysis of the blood-meal, which he also suggested was influenced by the development of intestinal bacterial symbionts. It has long been known that triatomine bugs have bacterial symbionts in their guts (Wigglesworth, 1936) which in the case of H. prolixus was named by Brecher and Wigglesworth (1944) as Acklimonyx rodhiki. However, the function of these symbionts is not as Dias (1934a) thought, to aid

digestion or hemolysis, but rather to provide missing nutritional factors which may be absent from the host's blood. Several studies have been carried out to determine specifically which factors the bacteria provide, with some emphasis on their role in synthesizing vitamin B factors although this is still a subject of recurring debate and further experiments (Goodchild, 1955; Baines, 1956; Harington, 1960; Lake and Friend, 1968; Auer, 1974). Phillips and Hartman (1967) found that the rate of reduction in the proportion of bugs infected at each larval stage was not linear and concluded that while the rate of digestion was probably the most important single factor influencing the development of trypanosomes in the gut of the bug, a complex of physiological processes was probably involved.

It is clear from the literature that variation exists both in the intensity of an infection of individual bugs and in the proportion of susceptible individuals in a population. This variation may be due to environmental factors affecting the bug and the parasite, but may also be affected by the bug genotype.

The possibility that the species of a bug may affect the vector-trypanosome relationship has been investigated by several workers. Dias (1940a), using a single Brazilian strain of *L. trumai*, obtained infection rates of 80-90% with Brazilian vectors of the disease (*T. infestans*, *P. megistus* and *T. cruzi*) but was only able to infect 56% of a group of *K. tritarsus* from Venezuela with this strain of trypanosome. Dias (1940b) tested the same colony of *P. prolixus* with a Venezuelan strain of *T. cruzi* and obtained the same infection rate (56%) but when *P. megistus* and *L. infestans* from Brazil were tested with the same Venezuelan strain of *T. cruzi* they showed a much reduced infection rate (40% and 36% respectively). Similarly, Floch and Lejolle

(1945) using local strains of T. cruzi in French Guiana found that 70% of K. pictipes and 85% of K. prolixus developed infections while only 31% of T. rubriventris could be infected, the degree of infection varying with their importance as vectors. Conversely, Condon and Viste (1957) found that T. dimidiata, the natural vector of T. cruzi in Costa Rica, with an infection rate of 76% was less susceptible than 4 exotic species T. phyllosoma (from Mexico) 81%, T. trinidadensis (from Chile) 86%, K. urulizus (from Salvador) 100% and K. laticornis (from Panama) 94%. Phillips and Bertran (1967) fed adult bugs of 4 species on rats with high levels of parasitaemia and found K. prolixus to be less susceptible (with infection rates 64-98%) than the other species T. dimidiata, T. protracta and T. muricola (infection rates of 89-100%). Little et al. (1961), using Trypanosoma brucei of T. cruzi found T. bahiensis, a species of trypanosome common in Mexico, to be more susceptible than T. brucei from Chile and concluded that inter-specific differences in bug physiology produced this variation between species. Schneider et al. (1973) fed 1 species of trypanosome larvae on 3 patients with chronic Chagas' disease in Argentina and T. infestans larvae gave a much higher infection rate (85%) than other species, for example T. pallidulus (1967). The higher infection rate of T. infestans being attributed to the fact that this is the most common vector in Argentina and therefore the strains of trypanosomes and the bug were co-adapted. It may be concluded from this work that variation in infection rates are related in part to the species of the bug which is in turn determined by its genotype.

The infection rates cited above demonstrate that there exists variation between individuals within a species which may be expressed in two ways, firstly, as a 'susceptibility rate' (the proportion of

susceptible individuals in a population expressed as a percentage) and secondly, as variation in 'intensity of infection' (the number of trypanosomes which an individual bug will support). Phillips and Bartram (1967) showed that individual bugs of 4 species were unable to support an infection despite heavy and repeated intakes of large numbers of parasites from host rats. This refractoriness they thought might have a genetic basis and by selecting refractory parents from a group of R. rattus with a susceptibility rate of 8% and inbreeding them, they produced offspring with a susceptibility rate of only 5% which suggested that susceptibility may be genetically controlled.

Few attempts have been made to quantify differences in the numbers of parasites supported by individual bugs. Wood (1960) counted the number of metacyclic and non-metacyclic trypanosomes in adult T. proutieri experimentally infected with T. evansi to determine the effect of temperature on the rate of trypanosome development in the bug. Wood (1960) constructed an infectivity index which compared the proportions of metacyclic to non-metacyclic trypanosomes per faecal droplet. Phillips and Bartram (1967) assigned an arbitrary numerical value to individual bugs using a scale from 1 to 6 by estimating the numbers of trypanosomes in the first faecal droplet released by an adult bug after feeding. Using this scale they observed differences in individuals of a group but not between the parental and F_1 generations of T. proutieri or Rattus norvegicus, although they did observe slight differences between species using this technique.

Wood (1960) noted a difference in intensity of infection between sexes of adult T. proutieri: he counted 1954 metacyclic trypanosomes per male (mean of 9 males) and 1821 per female (mean of 6 females). Phillips and Bartram (1967), however, found no significant difference

between sexes in any of the bug species they used, although they did note that females showed a slightly higher susceptibility rate (82%) than male H. hyalimorpha (64%) infected with the same strain of T. parvi. Likewise, Almeida et al. (1978) showed no difference in susceptibility between adult H. reuteri and found no difference in susceptibility rate between sexes.

It appears that only Phillips and Bertoni (1967) have compared the ability of different colonies of the same species of bug to transmit T. parvi: an inbred laboratory culture of H. hyalimorpha and a recently isolated field stock, showed no significant differences in susceptibility rates or in intensity of infection between the 2 populations studied. However, when they compared an inbred laboratory stock of H. infusum with field isolates of the same species, they found that the 3 wild stocks had an overall susceptibility rate of 100% while the laboratory stock had a slightly reduced rate of 98%. They also scored a small quantitative difference in intensity of infection of individuals between the 3 wild stocks but again this was not significant. It is not clear from this evidence if intensity of infection differs in individual bugs in a systematic fashion which could be related to genetic control, but it appears that such variation exists and may be due in part to the bug genotype.

Variation in the behaviour of different strains of T. parvi was first recognised by Brumpt (1913c) who noted that the virulence of two strains differed. Characteristics other than virulence (as measured by mortality) which have been reported to differ among strains are morphology, aspects of early growth, tissue affinity, course of parasitaemia, drug susceptibility, response to temperature, prepatent period and elimination of various antigens (Lindie, 1970). Variation in phenic characteristics

has been shown to affect the bug-trypanosome relationship. Phillips and Bartray (1967) noted a slight variation between strains in ability to infect bugs and this difference was constant for 4 species of bug. Sherlock and Almeida (1973) fed 10 species of triatomine bugs on dog, armadillo and mouse infected with 3 different strains of T. cruzi on dog, T. infestus and T. pallidus showed the highest susceptibility rates; on armadillo, T. infestus and T. brevipalpis had the highest rates, while on mouse, R. prolixus and T. trinidadensis had the highest rates. Means (1960) showed that when both American species of triatomine bug were infected with South American strains of T. cruzi a much lower density of metacyclic trypanosomes was found in the rectum than when the same species of bug were infected with a North American strain of T. cruzi and vice-versa. Reynolds (loc. cit.) concluded that strains of T. cruzi in nature are adapted to populations of triatomine bug from the same geographical region.

Various vertebrates have been shown to differ in their susceptibility to T. cruzi. For example, it has long been known that birds are not susceptible (Brumpt, 1914) and the guinea-pig is known to have poor susceptibility in the laboratory (Phillips, 1958), although it is said to be a natural host of the parasite (Means, 1960). It has also been shown that the resistance to infection increases with the age of the vertebrate host (Regusdanz, 1930; Dias, 1934; Kolodny, 1939a, b; Culbertson and Kesseler, 1942) and the severe forms of Chagas' disease in man are commonly encountered in infants. The strain of laboratory host is also said to influence its susceptibility suggesting genotype involvement. Fickel et al. (1960) showed that blood from mice interestingly susceptible to the virus homogenized they became. The sex of the host has also been examined for its effect on susceptibility and

Mauschke (1947) demonstrated that male mice were more susceptible than females of the same stock to 3 strains of T. cruzi. There is also evidence to suggest that environmental temperature affects the course of infection in the host; Kolodny (1939c, 1940) correlated variations in intensity of infection in experimental rats with seasonal temperature changes and Mauschke (1949) noted a temperature-dependent seasonal rhythm in the blood-trypanosome level of infected mice over a 27 month period. It has also been suggested that differences in behaviour within strains can be produced in the vertebrate host by altering the size of the inoculum given to the host animal. Phillips (1960a) investigated this by quantifying the parasitaemia of infected animals and found that the intensity of infection of 2 strains of T. cruzi measured in terms of parasite density in the blood and of virulence, varied with the number of parasites initially inoculated. Failure to inoculate sufficient parasites resulted in numerically attenuated infections with these 2 strains but in a third strain examined, the number of parasites inoculated proved immaterial. Mercedes (1967) and Mercedes and Hagstrom (1968), in experiments with mice and dogs, found that the pathogenicity of the 'Peru' strain of T. cruzi was related to the dose of infecting organisms and furthermore that the route of inoculation could influence the course of the disease. Mehlbein and Ormerod (1973) found great similarities between infections in mice both in intensity and mortality produced by trypanosome doses ranging over 4 orders of magnitude. Phillips (1964a) concluded that, given a sufficient number of parasites is inoculated to produce the maximum pathogenicity of which a strain is capable, the trypanosome will behave in a stable fashion and Mercedes (1967) found that mice given more than 1000 trypanosomes always died. It seems, therefore,

that the importance of the inoculating dose is still a matter for debate, although it is reasonable to suppose that dose and site of inoculum may have some effect on the course of an infection. It has also been suggested that the virulence of strains of T. cruzi could be lost over long periods of passage in laboratory animals (Galliard, 1952) but Hauschke (1949) found that passaging two strains of T. cruzi in mice for over 2 years increased their virulence. Phillips (1960a) quantitatively assessed a 'B.H.' strain infection of T. cruzi and found that it sustained its virulence unchanged for 2½ years in routine mouse passage.

From the work reviewed above it appears that several factors may affect the bug-parasite relationship and therefore, in order to study specifically the effect of bug genotype, factors other than this must be held constant if possible, including those affecting the bug (species, population, age, sex and external environment) and the trypanosome (strain, vertebrate host, host characteristics and inoculum).

The present work describes experiments designed to breed susceptible and refractory populations of K. jellisoni. A pure breeding susceptible population would have great value in serodiagnosis.

MATERIALS AND METHODS

Materials

One strain of colonized K. jellisoni, 2 strains of Trypanosoma cruzi and T.O. strains were as vertebrate hosts. K. jellisoni. A single species of triatomine, R. prolixus (Lai, 1959), was used in this experiment and was chosen because of its relatively short life-cycle (egg to adult taking about 6 months at 28°C (cardinal

and Maddrell, 1972)) and because it requires only one blood-meal between moults (Wigglesworth, 1934). The L,S,H, and T,M, colony of R. prolixus is now a highly adapted laboratory strain, an indication of its fitness is that in 1930 the fertility of the colony was estimated to be about 82% (Muxton, 1930) and is now about 93% (see Part Four).

1. Strain 6. Two strains of T. cruzi were used in this experiment and were chosen for their well-documented, and differing characteristics.

1. Peru Strain. A strain of high virulence, isolated in 1948 by Dr. F. Naquina in Aiquepa, Peru. It was given that year to Dr. F. Goble (Russek-Weig and Goble, 1956) who in turn gave it to Dr. P.D. Marsden of L,S,H, and T,M. (Marsden, 1957). The strain was subsequently passaged over 160 times in mice but at passage 155 it was stabilized by standard methods of cryopreservation as reference code IATP 693 (for details of this procedure see Marsden et al., 1973). This stabilate was then used for the present experiment, and is herein referred to as the Peru strain. This strain has a pre-patent period of 8-14 days in mice and is lethal at 25-30 days in inocula containing more than 100 organisms (Matsubata and Ormerod, 1971).

2. Strain 7. A low virulence strain of T. cruzi, isolated in 1965 from an acute case of Chagas' disease in man in Sepilip, Brazil, by Dr. P.D. Marsden and brought to the L,S,H, and T,M. colonies in 1966. The strain was then passaged in mice 12 times, stabilized as IATP 518 and 519, passaged once more in mouse and stabilized as reference code IATP 692. This stabilate was used in this work. Strain 7 is not lethal to mice and they may recover from inocula of over $\frac{1}{2}$ million trypanosomes. The pre-patent period in mice is 8 days and after 30 days parasitemia declines and the mice recover

(Kettleridge, personal communication).

Test animals used. Six to eight week old male mice of the Jackson bred T.O. (Theiler's Original) strain weighing 24-30 g were used throughout and maintained in one room at 24°C.

Methods

The protocol adopted for this experiment to ensure the synchrony of infected mice with bugs of the correct age is shown in Fig. 3.2 and was adhered to over successive generations of bugs.

Treatment of mice. Stabilates of both strains of L. smithi were inoculated into mice and the infection followed by examination of tail blood one week after inoculation and subsequent bleedings. The procedures followed in handling infected material are detailed in Lumsden et al., 1973). Mice with a parasitaemia of 1-5 trypanosomes per field (estimated by scoring 100 fields, X 450) were bled from the heart. 1 ml of infected blood was diluted with 5 ml heparinized salt solution (Oxoid, England) and 0.1 ml of this diluted blood was injected intra-peritoneally (i.p.) into new experimental mice. This standardized system, although subject to variation in the numbers of trypanosomes ingested, did produce very reproducible patterns of infection in the experimental mice as measured by tail bleeding. Mice infected with the Peru strain regularly developed parasitaemia of 1-5 trypanosomes per field (100 fields, X 450) by examination of tail blood 3 days post-inoculation. Mice infected with Strain 7 developed a parasitaemia of approximately 1 trypanosome per field 3 days after inoculation by the same method of examination. These results were consistent throughout the experiment. Although a series of previous work suggested that the number of trypanosomes ingested by a bug was not a critical factor in determining the level of infection developed by the bug, this

standardized method of mouse inoculation was an attempt to ensure that bugs were ingesting approximately the same numbers of *Trypanosoma* in terms of orders of magnitude. With such large numbers of mice involved in this experiment, it would have been difficult to ensure that each mouse had exactly the same level of parasitaemia.

Treatment of bugs. *R. prolixus* were kept in an incubator at a constant temperature of 28°C and R.H. of approximately 70% and contained in 2" x 11" flat-bottomed glass tubes, sealed with fine mesh nylon gauze held in place by adhesive tape. Since the care and feeding of larval stages may influence the susceptibility to disease of adult insects (Burt, 1946), great care was taken to provide optimal and identical rearing conditions for the larval stages of experimental bugs. Larvae were reared in tubes with only a few larvae per tube to preclude competition when feeding, and the tubes of larvae were maintained in incubators under the same conditions of temperature and humidity described above. Larval blood-meals were given on uninfected mice in the same way as described for infecting feeds. To ensure that the experimental bugs were all of the same age when given their infecting feed, they were all fed simultaneously as fourth instar larvae, allowed to moult to fifth instar larvae and, as fifths were given the infecting blood-meal exactly 28 days after this fourth instar blood-meal. Fifth instar larvae were chosen for the infecting feed because they take more blood than any other stage of *R. prolixus*; a feed of 200-300 mg blood is usual for fifth instar larvae (Wigglesworth, 1934) which guaranteed an adequate intake of *trypanosomes*.

Infecting feed. For the infecting blood-meal the fifth instar larvae were placed in pairs in tubes, filter paper having been placed in the tubes to assist their stance while feeding and to absorb their faeces. The parasitaemia of each infected mouse was first estimated by

standardized method of mouse inoculation was an attempt to ensure that bugs were ingesting approximately the same numbers of trypanosomes in terms of orders of magnitude. With such large numbers of mice involved in this experiment, it would have been difficult to ensure that each mouse had exactly the same level of parasitemia.

Treatment of bugs. *R. prolixus* were kept in an incubator at a constant temperature of 28°C and R.H. of approximately 75% and contained in 2" x 1½" flat-bottomed glass tubes, sealed with fine mesh nylon gauze held in place by adhesive tape. Since the care and feeding of larval stages may influence the susceptibility to disease of adult insects (Hurst, 1947), great care was taken to provide optimal and consistent rearing conditions for the larval stages of experimental bugs. Larvae were reared in tubes with only a few larvae per tube to preclude competition when feeding, and new tubes of larvae were substituted in incubators under the same conditions of temperature and humidity described above. Larval bloodmeals were given on scheduled mice in the same way as described for infecting adults. To ensure that the experimental bugs were all of the same age when given their infecting feed, they were all fed simultaneously as fourth instar larvae, allowed to moult to fifth instar larvae and, as fifths were given the infecting bloodmeal exactly 28 days after their fourth instar bloodmeal. Fifth instar larvae were chosen for the infecting feed because they take more blood than any other stage of *R. prolixus*; a feed of 210-300 mg blood is usual for fifth instar larvae (Wigglesworth, 1934) which guaranteed an adequate intake of trypanosomes.

Infecting feed. For the infecting bloodmeal five fifth instar larvae were placed in pairs in tubes. Filter paper having been placed in the tubes to assist larval moult while feeding and to absorb blood excreta. The parasitemia of each infected mouse was first estimated by

examination of tail blood on the day of feeding and mice were selected for the feeds using the criteria described above. Mice used for feeding bugs were anaesthetised with 0.2 ml of a 20% solution of 'Nembutal' (Pentobarbitone sodium B. vet. Co.), administered i.p. and, after sedation, they were placed on top of the tubes containing the bugs so that each mouse covered 2 tubes and thereby fed 4 fifth instar larvae. The tubes containing the bugs were placed upright randomly in a polystyrene box prior to the feed, the lid being placed on the box after the mice had been laid on the tubes, to provide insulation. Since *N. violaceus* is attracted to its host from short range mainly by the warm air diffusing from it (Wigglesworth and Gillett, 1934), insulating the mice in this way prevented the fall in body temperature normally associated with this anaesthesia. After the bugs had fed, they were returned in their tubes to the incubator at 28°C. The mice were killed with ether, before they recovered from the anaesthesia, and incinerated.

One week after the infecting feed, and prior to the moult to the adult, the bugs were placed individually in clean glass tubes which prevented any unplanned matings. Following the moult the adult bugs were placed in clean tubes which, instead of filter paper, contained a strip of nylon mesh which enabled the bugs to climb up the tubes to feed on a clean mouse when offered it, due course, but would not absorb any of the faeces which would be required for examination for trypanosomes. At this stage also, bugs were sexed and a code number was written on the side of each tube for identification.

Measurement of bug lifespan.

1. Time of examination. Mackell (1964) examined bugs for xenodiagnostics 40-60 days after the infecting feed and Freitas (1950)

examined bug, 30, 60 and 90 days after feeding and found that the group of bugs examined at 60 days had a significantly higher infection rate than the other 2 groups. Patterson and Miles (1973) found that the intensity of infection in the rectum of H. polydora fed on rhesus monkeys infected with T. evansi did not remain constant and was highest 25 days after the blood-meal. In the present experiment all adult bugs were examined exactly 35 days after their infecting feed as fifth instar larvae (Patterson and Miles, loc. cit. have shown that metamorphosis and ecdysis of fifth instar larvae to adults have no effect on the intensity of infection in the gut of the bugs).

2. Method of examination of faeces. Since the bugs were required to mate to produce progeny for these genetic studies, manual extraction of rectal contents not squeezing the abdomen could be employed for faecal examination for trypanosomes, and therefore bugs were fed on uninfected mice and were allowed to defaecate naturally during and after this 'clean feed'. For this 'clean feed' the same apparatus was used as for the infecting feed. The anaesthetized, clean mice were placed on top of the bug tubes, randomly arranged in a polystyrene box, so that each mouse fed 2 bugs. But, for this meal, 1 ml Hank's solution was injected to the bottom of each bug tube using sterile graduated syringes, and the bug faeces, as they were excreted, dropped from the nylon mesh in the tube into the Hank's solution in which desiccation of the trypanosomes was prevented. Any faecal material remaining on the nylon mesh strip was mixed with the Hank's solution before sampling. Wigglesworth (1931) showed that H. polydora voids the residue of its previous meal immediately after feeding, then a few minutes later excretes a cloudy, watery fluid and for the next 3-4 hours passes clear, colourless, urine at intervals, although the

rate of excretion declines rapidly 1-2 hours after feeding. Since the faeces of many bugs often had to be separately examined on the same day, it was not possible to control strictly the time interval between feeding a bug and examination of faeces, but when numbers were large on the same day, the bug tubes were grouped randomly and groups fed at conveniently spaced intervals, so keeping to a minimum variation in the time between feeding, excretion and examination of the faeces. Using this system the faeces of most bugs were produced and available for microscopic examination between one and two hours after feeding. This seemed to be a reasonable and practicable procedure. The clean mice were killed immediately after use, before their bedding bags had further chance to exclude the possibility of faecal infection.

3. Scoring of faecal infection. The number of parasites excreted by each bug was estimated by sampling the Marks solution after it had been well mixed with the faeces. Sampling was carried out using calibrated glass micropipettes (Barnes Scientific Co., U.S.A.), a clean pipette being used for each bug's faeces. A micropipette was pushed through the gauze covering the top of a tube containing a bug and its faeces in Marks solution, and allowed to fill by capillary action. There 20 samples were then expelled on to glass microscope slides using a rubber bulb, No. 1 coverslips (10 x 25 mm) (Chance Wrought Ltd., England) were placed on the droplets and the slides were marked with code numbers corresponding to those on the tubes containing the bugs.

Scale of examination.

It would perhaps have been more accurate to have used a haemocytometer to estimate the numbers of trypaenosomes in the samples but

this would have involved too much time per insect, making the assessment of infection in a large number of bugs impossible. Moreover, haemocytometers have an inherent error related to mathematical considerations of the patterns in which organisms are expected to occur in the squares of the apparatus (Dacie and Lewis, 1963). The method chosen was to count 100 fields (X 450, using phase contrast microscopy) of each sample, working in a systematic way across the coverslip in stepwise increments. Only one sample from each bug was examined except for bugs of the F_3 generation infected with T. brucei strain F_3 , when 2 samples were taken from each bug to obtain a measure of the error variation between samples. Working counted the number of trypanosomes in 100 fields; this number was divided by the bug as a 'count'. It must be noted that all forms of trypanosome encountered in each sample were counted; previous workers have sometimes distinguished between the transmissible, infective metacyclic forms and other forms (amastigotes and epimastigotes, Mahabadi and Denny, 1973) which are sometimes found in the faeces but since the production of trypanosomes of whatever stage of development is evidence that the vector is supporting the development of the parasite, the counting of all forms, rather than time-consuming differentiation of different forms, seemed to be both a practicable and adequate measure of vector infectivity. Moreover, for the purpose of serodiagnosis, the evidence of trypanosomes in the faeces, of whatever developmental stages, is proof that the original donor patient, or animal, is infected.

If no parasites were seen in a faecal sample, the bug was kept and given a second 'clean feed' 14 days later when, if it proved negative again, a sample of the diluted faeces was injected i.p. into a clean mouse. The tail blood of the mouse was examined for trypanosomes

14 and 21 days later. If the bug showed no evidence of parasites on all three tests it was assumed to be refractory to infection, and assigned a score of zero. If a bug was negative at the first test but proved positive at a later test it was assigned an arbitrary score of one, since the second defaecated sample or, the final assay (the recipient mouse parasitaemia) could not be quantitatively related to the first defaecated sample scores of other bugs in the experiment.

Methods of selection of bugs for breeding

The parental generations of *M. tritoxus* were selected randomly from the stock colony as fourth instar larvae and, after feeding and moulting to the fifth instar, 2 groups of bugs, each consisting of 50 male larvae and 50 female larvae were given an infecting blood-meal on mixed as described above. These 2 groups of bugs were 1-100 designated 'Peru' and 'Hawaii' according to the place of the original origin. Both groups were allowed to moult to adult, then fed on clean mice, and their faeces scored as described above. Bugs which did not orally engorge at an infecting meal were rejected not only in these parental groups but also in subsequent generations.

Selection of bugs as parents for the next generation was made on the basis of individual faecal scores for trypanosomes. Selected males from each of the parental groups were mated with several females from the same group and 3 blood-meals given to each pair at 14 day intervals. The eggs laid by each female over the 3 feeds were pooled, each group of eggs representing an F_1 family. On emergence, the F_1 first instar larvae were fed on clean mice and reared to the fifth instar when they were given an infecting feed and subsequently scored, now adults on the basis of faecal infection. Single pair within assortative matings were made from these F_1 adults using individual

scores as the basis for selection. This process of infecting feed, followed by scoring, selection, and assortative mating, was repeated to produce an F_2 generation from both 'Peru' and 'Strain 7' bugs. However, only the 'Strain 7' group was reared to the F_3 generation, the 'Peru' experiment being discontinued at the F_2 generation.

Methods of data analysis

1. Analysis by class

For a major gene interpretation, infected bugs fall into 2 obvious discrete classes, (a) refractory - bugs with no trypanosomes in their faeces, and (b) susceptible - bugs with trypanosomes in their faeces. It is not essential in analysing such data to give a numerical definition of the individuals within the classes defined; it is sufficient that the classes are clearly distinct producing a discontinuous distribution so that analysis is based simply on the number of individuals in each class. The present experiment examined the genetic structure of 2 populations of bugs, Strain 1 and Peru, through generations of selection in order to examine any changes in proportion of refractory to susceptible bugs which would allow for a Mendelian interpretation of the control of susceptibility.

2. Analysis by intensity of infection

Analysis of continuous variation presents different problems, for each observation is important and reflects the expression of the character. The number of trypanosomes excreted by a bug cannot be continuous score ratio, the fraction of trypanosomes in the excreta and it is therefore essential to treat the scores as if such minor discontinuities did not exist, that is as 'quasi continuous' variation (Falconer, 1953). In quantitating and analysing intensity of infection it is therefore necessary to replace frequencies by means

and variances. For the application of most statistical methods to a set of data, its frequency distribution should be normal, but the dispersal pattern of trypanosomes is not normal since they tend to be clumped or aggregated in space. This distribution, which is described as 'contagious' (Southwood, 1966) is non-random and in order to normalize it, the data must be transformed; that is, the actual numbers must be replaced by a function whose distribution is such that it normalizes the data. The appropriate transformation for trypanosome population estimates has been arrived at empirically and it has been shown that a logarithmic transformation is most suitable (Harden et al., 1971). Accordingly, in assessing the intensity of infection of bugs in the present work all scores were transformed logarithmically, and, in order to overcome difficulties with zero counts, a constant was added to the original scores so that all scores were transformed to a $\log(n + 1)$ scale (where n represents the score) before analysis.

A quantitative analysis was carried out on the transformed scores of the bugs in the present selection experiment to allow for the possible polygenic control of susceptibility.

RESULTS

Numbers of bugs selected

Larv. Of the 100 fifth instar larval bugs initially infected with Peru strain *T. cruzi*, 81 fed fully, moulted successfully and were scored as adults, 41 of which were male and 40 were female. Eleven of these 81 bugs had a cast vomit. During collection of all faeces of their faeces, and of these 11 refractory bugs, 6 (3 males and 4 females) survived and were used for mating. Five arrays of

half-sib families were established, being 4 males each mated to 4 females. Some matings were unsuccessful and only 10 of the 16 matings remained. In addition, 2 further matings were established using randomly chosen parents, giving rise to 12 F_1 families in all (see Table 1.1).

These F_1 families were reared, scored as adults and 19 full-sib matings were selected from among 8 of the F_1 families. Fifteen of these matings were selected for high score and 4 for low score. The offspring of these full-sib assortative matings were reared to adults, infected and scored. The Peru experiment was not taken further than this F_2 generation.

Strain 2. From a large population of 150 first instar larvae infected with Strain 2 larvae, 77 (41 males and 36 females) were reared successfully to the adult stage and scored. Of 77 bugs, 8 had scores of zero and were found to be refractory on all tests of their parents. Of 14 males, 2 females of these 8 scored just one used for mating. The design of these matings was similar to that for the Peru bugs consisting of 4 males each of which was mated to 4 females (see Table 1.1). Eleven of these 12 F_1 bugs produced viable F_1 families which were reared, infected and scored as adults. On the basis of these F_1 scores, 14 full-sib assortative matings were established together with 2 unrelated crosses from among 9 of the 11 F_1 families. These 16 F_2 families were reared and scored as adults and from 9 of these families further selections were made and 17 full-sib matings set up. These 17 selected pairs were divided into high and low lines. Four pairs were assortatively mated for high score and from the same F_2 families 4 pairs with low scores were mated to provide reverse selection controls. Seven pairs were mated assortatively for low score and 4 reverse selections were also

established from these low line families, but only 2 of these control matings were fertile.

The progeny of these 17 full-sib F_2 matings was reared, infected and scored as F_3 adults. These F_3 individuals were scored using 2 faecal samples being examined from each bug to increase the precision of the F_3 statistics.

Analysis by class

Peru. The parental matings made from the Peru population are shown in Table 3.1 in which bug scores are presented untransformed. F_1 families were coded according to the number given to their female parent. F_1 families 73 and 61 were produced from crosses in which both the parents were refractory (i.e. score Peru). Family 73 was bred to produce F_2 families 17 and 18 using only refractory parents while family 61 was similarly bred to produce F_2 family 16. The results of this breeding experiment over 2 generations of selection are shown in Table 3.2 where results are expressed in terms of the proportions of refractory to susceptible bugs in the families of each generation raised.

Strain 7. The 4 groups of parental matings selected from the base population of M. trilineatus infected with L. peru Strain 7 are shown in Table 3.3.

F_1 families 93 and 94 were produced from parents with mean scores. F_1 family 93 was bred successfully to F_3 family 2 using only refractory parents, and family 94 was bred to F_3 family 12 using parents with high scores in selecting for susceptibility. The results of selection and breeding for susceptible and refractory bugs are shown in Table 3.4. Only those F_1 and F_2 families which provided parents for subsequent generations are included in Table 3.4.

Table 3.1. Twelve matings from base population E. prolium infected with Peru strain T. cruzi showing code numbers and scores (untransformed).

Array 1			Array 2			Array 3			Array 4			Random Mating					
Sex	Bug No.	Score	Sex	Bug No.	Score	Sex	Bug No.	Score	Sex	Bug No.	Score	Sex	Bug No.	Score	Sex	Bug No.	Score
♂	37	11	♂	52	0	♂	1	0	♂	40	4	♂	25	4	♂	66	3
Q	21	0	Q	73	0	Q	61	0	Q	20	0	Q	43	4	Q	63	1
Q	48	3	Q	55	3	Q	58	5	Q	56	4						
Q	6	3	Q	2	9												

* No. of trypanosomes/100 fields

Table 3.3. Frequencies of refractory bugs on the V_1 and V_2 generations of *S. aureus* Peru strain infected *S. proluxus*. Scores are shown untransformed.

Family number	Parental score		T	n	with zero score	Family number	Parental score		T	n	% with zero score	Direction of selection
	P	Q					P	Q				
43	4	4	15		0	3	4 3 2	24 10 5	18 8 15	1	5.6 0 6.7	S S S
63	3		28	4	14.3	4	20	5	15	2	13.3	S
55	0	3	36	1	2.8	6 7 8	67 25 96	29 24 27	15 13 12	1	6.7 7.7 8.3	S S S
56	4	4	24	2	8.3	9 10	6 6	23 3	19 19	2 1	10.5 5.3	S S
2	0	9	37	1	2.7	11 12 13 18	19 15 74 141	20 25 24 11	20 11 19 20	2 3	0 0 10.5 15.0	S S S S
Total									204	15	7.3	

Table 3.2. (Continued)

F ₁						F ₂						Direction of selection
Family number	Parental score ♂ ♀		T	n	% with zero score	Family number	Parental score ♂ ♀		T	n	% with zero	
61	0	0	43	5	11.6	16	0	0	14	1	7.1	R
73	0	0	14	4	14.3	18	0	0	16	3	18.8	R
6	11	3	9	2	22.2	19	0	0	16	2	12.5	R
21 x 48	11	3	23	1	5.7 4.4	20	0	0	16	2	12.5	R
2 x 55	0	9 3	37 36	1	2.7 2.8	21	0	0	20	0	0	R
Total									99	11	11.1	

T = Total No. bugs scored

n = No. of refractory bugs (with zero score)

R = Selection for refractoriness

R = Selection for susceptibility

Table 1.3. Slaves caught from nine populations of wildfish collected with French 7
G. trawl, showing code numbers and scores (untransformed)

Array 1			Array 2			Array 3			Array 4		
Sex	Bug No.	Score	Sex	Bug No.	Score	Sex	Bug No.	Score	Sex	Bug No.	Score
♂	51	0	♂	42	0	♂	17		♂	15	0
Q	50	3	Q	93	0	Q	94	0	Q	70	7
Q	69	30	Q	73	7	Q	58	9	Q	76	14
						Q	46	17	Q	66	20
						Q	55	61			

No. of transformation factor

Table 1.4. Frequencies of refractory bugs in the F_1 , F_2 and F_3 generations of La strain Strain T infected S. anitima.

Numbers are given untransformed.

F ₁						F ₂						F ₃						
Family number	Parental score		T	n	% with zero score	Family number	Parental score		T	n	% with zero score	Family number	Parental score		T	n	% with zero score	Direction of selection
	♂	♀					♂	♀					♂	♀				
93	0	0	22	5	22.7	10	0	0	17	2	11.8		138	10	23	6	25.1	Ra
												0			24	5	20.8	R
70	0	7	22	10	45.5	11	0		19	7	36.8	4			31	1	3.2	R
												5	0		13	0	0	R
73	1	7	26	11	55.0	13	0	0	21	3	14.3	7	0	0	37	2	5.4	R
69	0	30	22	2	9.1	15	0	0	20	2	10.0	8	86	42	13	1	7.7	Ra
												9	0	9	18	1	5.6	R
73	1	7	20	11	55.0	14	0	0	21	3	14.3	10			24	10	41.7	R
												11	0	0	30	14	46.7	R
Total													213		40	18.7		

..... Continued

Table 3.4. (Continued)

F ₁						F ₂						F ₃						Direction of selection
Family number	Parental score		T	n	% with zero score	Family number	Parental score		T	n	% with zero score	Family number	Parental score		T	n	% with zero score	
	♂	♀					♂	♀					♂	♀				
50	0	3	36	3	8.3	2	156	33	15	0	0	12 13	38 2	14 6	35 8	2 0	5.7 0	S Sr
94	0	0	36	0	0	5	118	103	21	1	4.8	14 15	114 4	20 0	33 24	0 1	0 4.2	S Sr
70	0	7	22	10	45.5	8	184		17	1	5.9	16	370	30	27	0	0	S
46	1	17	34	3	8.8							17	2	0	24	0	0	Sr
58	1	9	26	1	3.8	3	286	256	26	0	0	18 19	355 6	88 4	24 10	1 1	4.2 10.2	S Sr
Total														5		2.7		

T = Total No. bugs scored

n = No. of refractory bugs (with zero score)

Rs = Reverse selection from refractory

Sr = Reverse selection from susceptible

Differences between resistant populations

The proportions of refractory bugs in the 2 base populations were compared by a χ^2 test, the results of which are shown in Table 3.5.

Table 3.5. χ^2 test for difference in proportions of refractory bugs between 2 base populations of *M. gutierrezii* infected with *T. cruzi* Peru strain and Strain 7.

Bug population	T	n	mean % refractory	χ^2	P
Peru	81	11	13.6	0.008	0.92
Strain 7	77	8	10.4		

Legend to Table 3.5

- n.s. = Probability is non-significant
- + = Probability = 0.05 - 0.10
- * = Probability = 0.01 - 0.05
- ** = Probability = 0.001 - 0.01
- *** = Probability = <0.001
- P = Probability

The results show that there was no significant difference in the proportion of refractory bugs between the 2 base populations of *M. gutierrezii* infected with the 2 different strains of *T. cruzi*.

Differences between selection lines

The effect of selection for susceptibility and refractoriness on the proportions of the 2 types after successive generations of selection

was compared for both groups of bugs infected with different strains of T. cruzi:

(a) Peru. The proportions of refractory bugs in 19 F_2 families of 2 selection lines, were compared by a χ^2 test (Table 3.6.).

Table 3.6. χ^2 test for differences in proportion of refractory bugs between 2 selection lines from 19 F_2 families of R. prolixus infected with Peru strain T. cruzi.

Selection line	T	n	%	χ^2	P
Susceptible families	204	15	7.3	0.77	N.S.
Refractory families	99	11	11.1		

The results show that selection for refractoriness and susceptibility over 2 generations of assortative mating did not significantly alter the proportions of the 2 classes of bug in the F_2 generation of Peru strain infected R. prolixus.

(b) Strain 7. The proportion of refractory bugs in 9 F_2 families infected with T. cruzi Strain 7 were compared by a χ^2 test (Table 3.7.).

Table 3.7. χ^2 test for differences in proportions of refractory bugs between 9 F_2 families from 2 selection lines of R. prolixus infected with T. cruzi Strain 7.

Selection line	T	n	%	χ^2	P
Susceptible families	79	2	2.5	18.7	<0.001***
Refractory families	98	17	17.4		

The results show that for B. prolixus infected with T. cruzi Strain 7, after 2 generations of assortative mating there was a highly significant difference in the proportions of refractory bugs between families selected for susceptibility and those selected for refractoriness.

A similar analysis was carried out on the proportions of refractory bugs found in F_3 families infected with T. cruzi Strain 7 (Table 3.8).

Table 3.8. χ^2 test for differences in proportions of refractory bugs between 17 F_3 families of two selection lines of B. prolixus infected with T. cruzi Strain 7.

Selection line	T	n	%	χ^2	P
Susceptible families	.185	5	2.7	19.2	<.001***
Refractory families	.213	40	18.7		

The results show that the highly significant differences detected in generation F_2 were reproduced in generation F_3 in which highly significant differences in the proportions of refractory bugs between selection lines were found. These significant differences can only have resulted as a consequence of the selection pressure applied to the original population.

Differences between families within selection lines

The proportions of refractory to susceptible bugs within families are shown in Tables 3.2 and 3.4 together with the direction of selection applied to each family. The proportions of refractory bugs should have

increased in those families deliberately selected for refractoriness, and a test for linear trends in proportion was applied to selected families to detect any consistent change over generations. Three F_2 families from the Peru group, numbers 17, 18 and 19 were chosen since they were bred from parents and grandparents which were phenotypically refractory. The results of this analysis are shown in Table 3.9. These analyses were carried out following the method of Armitage (1955).

Table 3.9. Test for linear trend in the ratio of proportions of refractory bugs in 3 families of *E. coli* selected for refractoriness to T₁ using back crosses using P , F_1 and F_2 data

F_2 family number	\hat{b}	S.E.	χ^2/df	P
17	0.0090	0.0371	0.06	N.S.
18	0.0117	0.0377	0.10	N.S.
19	0.0027	0.0404	0.004	N.S.

\hat{b} = Slope of regression line

S.E. = Standard error \hat{b}

The results show that there was no significant trend towards an increase in the proportions of refractory bugs in these 3 families over 3 generations. Despite the inbreeding of refractory parents, F_2 family 16 which was bred from refractory parents and grandparents, in fact,

Table 1.18. Tests for linear trends in proportions of refractory R. prolixus in 9 selection lines selected for refractoriness to T. cruzi Strain 7 using P , F_1 , F_2 and F_3 data.

P_1 Family No.	b	S.E.	χ^2 , 3df	P
I	0.0321	0.0228	1.9846	N.S.
II	0.0099	0.0200	0.2475	N.S.
III	0.0011	0.0230	0.0019	N.S.
IV	0.0275	0.0294	0.8759	N.S.
V	-0.0175	0.0209	0.3523	N.S.
VI	-0.0056	0.0229	0.0587	N.S.
VII	-0.0103	0.0210	0.2376	N.S.
VIII	0.0563	0.0244	5.3126	N.S.
IX	0.0636	0.0233	7.4348	0.05 - 0.1
Total	-0.0068	0.0137	0.2393	N.S.

b = slope of regression line

S.E. = standard error b

showed a decreased proportion of refractory bugs, having only 7% while its parents, which were both refractory, came from an F_1 family (No. 61) with 11% refractory bugs.

Similar analyses were carried out using data from the F , F_1 , F_2 and F_3 generation families infected with T. cruzi Strain 7, which were selected for resistance to it. The results of these analyses are shown in Table 3.10. The results show that there was no highly significant trends towards an increase in the proportions of refractory bugs in any of these families from the parental to the F_3 generation. The most surprising result in this connection is the F_3 family which had parents, grandparents and great-grandparents all phenotypically refractory and yet showed no trend towards an increasing proportion of refractory members, having 20% refractory members compared with its parental family proportion of 11% and its grand-parental family with 22% refractory. Furthermore, pooling the data from all these families failed to show any significant linear trend towards increasing proportions of refractory bugs; indeed the pooled data produced a negative slope indicating a slight overall decrease in the proportions of refractory bugs.

A similar analysis was carried out using those families infected with T. cruzi Strain 7 which had been selected for susceptibility (Table 3.11).

Table 3.11. Test for linear trend in proportions of refractory bugs in 8 F_3 families selected for susceptibility to infection with T. cruzi Strain 7 using F , F_1 , F_2 and F_3 data.

F_3 family numbers	\hat{b}	S.E. \hat{b}	χ^2	P
12 - 19 pooled	0.0008	0.0092	9.76	0.01 - 0.05*

The results show that in these 8 families there was a significant linear trend downwards in the proportion of refractory bugs from the parental generation to the F_3 , $t = -0.71$ which indicates that selection for susceptibility in these families had indeed steadily increased the proportion of susceptible bugs.

It is difficult to give an explanation for the apparently conflicting results presented above. For example, after 3 generations of assortative mating, the Peru strain F_2 family with the greatest proportion of refractory bugs (F_2 family 18) did not have a significantly greater proportion of refractory members (18.6%) than the original parental generation (15.6%) (corrected $t = 1.04$, $P = N.S.$). Similarly, in the families selected for susceptibility, there was no significant overall increase in the proportions of susceptible bugs (84.6% to 85.7%, $X^2 = 0.6$, $P = N.S.$).

The F_3 families bred for refractoriness to F_2 family 7 had a mean proportion of 18.7% refractory members which was not significantly different from the parental generation level of 15.6% ($X^2 = 1.84$, $P = N.S.$). Moreover, the F_3 families with the greatest proportions of refractory bugs (>40%) were F_2 families 10 and 11 which were bred initially from F_1 family 7, the parents of which were not refractory (see Table 1).

The F_3 families selected for susceptibility did however show a significant increase in the proportion of susceptible members compared with the base population (from 85.6% to 87.3%, $t = 4.3$, $P = 0.02 = 2.04\%$) and this increase was fairly constant in all families. The results have not, however, demonstrated any radical changes in the proportions of either susceptible or refractory bugs as a result of selection which, assuming a major gene controlling factor, would have been expected after 3 generations of selection and self-mating.

Quantitative analysis

An alternative genetic analysis was attempted using the scores of individual bugs (i.e. intensity of infection) scores in the infection experiments described above. Instead of assigning each bug to a class (i.e. positive or negative for T. brucei) on the basis of its score, the actual score (number of trypanosomes per blood smear) for each bug was transformed and then used as raw data in computing first and second degree statistics for families and selection lines. The transformed data are presented as family and generation means in Tables 3.12 - 3.26. These data are based on scores from a total of 1725 K. burleti adults. The family mean scores, untransformed and transformed scores of selected parents have already been given in Tables 3.1 - 3.4 together with their selecting lines. The present analysis, selection for susceptibility implies low scores, and selection for susceptibility implies selection for high scores.

Differences between strains of T. brucei

The mean scores presented in Tables 3.12 - 3.26 show that intensity of infection was strikingly different in the bugs infected with Peru strain from the bugs infected with Strain 7 T. brucei.

Differences between the 2 parental generations were compared by analysis of variance of scores of bugs infected with Peru and Strain 7, analyses of male and female scores being carried out separately as well as together, and the results are shown in Tables 3.29, 3.30 and 3.31.

Table 3.12. Transformed data for Peru parental generation R. prolixus

Sex	n	\bar{x}	S.E.	S.D.
♂	41	0.4797	0.0408	0.2609
♀	40	0.5005	0.0428	0.2706
♂ + ♀	81	0.4900	0.0294	0.2643

Key to annotation for data Tables 3.12 - 3.28

n = Number of bugs tested

\bar{x} = Mean score

S.E. = Standard error of mean

S.D. = Standard deviation

Table 3.13. Transformed data for Peru F_1 generation male *B. prolixum*

Family number	n	\bar{x}	S.E.	S.D.
21	20	0.6021	0.0778	0.3478
48	10	0.4970	0.0710	0.2246
6	5	0.7360	0.2556	0.5716
73	15	0.8915	0.1133	0.4389
55	17	1.1400	0.1281	0.5281
2	23	1.1525	0.1200	0.5752
61	15	0.9130	0.1664	0.6443
58	5	1.0673	0.2648	0.5922
20	8	0.7456	0.1045	0.2954
56	12	0.6966	0.0704	0.2433
43	3	0.5927	0.0642	0.1112
63	15	0.5488	0.1125	0.4358

Table 3.1d. Transformed data for Paru F, generation female *S. polixus*

Family number	n	\bar{x}	s^2	S.D.
21	13	0.6119	0.1036	0.4012
48	13	0.5841	0.1035	0.3731
6	4	0.6851	0.3230	0.6460
73	18	0.4493	0.0777	0.3298
55	19	0.8620	0.1054	0.4595
2	14	1.0804	0.0983	0.3679
61	27	0.6396	0.0825	0.4288
58	3	0.6514	0.1901	0.3292
20	9	0.8412	0.1731	0.5193
56	12	0.5054	0.1161	0.4022
43	12	0.6899	0.1015	0.2517
63	11	0.5663	0.1212	0.4369

Table 3.15. Transformed data for Peru F_1 generation families
(male and female) of *B. prolixus*

Family number	n	\bar{x}	S.E.	S.D.
21	35	0.6063	0.0619	0.3659
48	23	0.5462	0.0654	0.3138
6	9	0.7134	0.1887	0.5662
73	33	0.6503	0.0763	0.4382
55	36	0.9933	0.0843	0.5054
2	37	1.1253	0.0826	0.5024
61	42	0.7372	0.0811	0.5253
58	8	0.9114	0.1863	0.5269
20	17	0.7962	0.1016	0.4189
56	24	0.6010	0.0693	0.3400
43	15	0.6795	0.0819	0.3172
63	28	0.5569	0.0809	0.4282

Table 3.16. Transformed data for Pam: F_1 generation male P. prolixus

Family number	n	\bar{x}	S.E.	S.D.
1	18	0.7169	0.1321	0.4178
2	4	0.7107	0.2747	0.5493
3	9	0.7584	0.1803	0.5805
4	4	0.9619	0.1772	0.4688
5	7	0.8219	0.2158	0.5710
7	8	0.7556	0.1712	0.4841
8	9	1.1554	0.2319	0.5185
9	13	0.6616	0.0758	0.2734
10	10	0.7816	0.0524	0.1657
11	11	1.1488	0.0828	0.2986
12	9	0.7487	0.1095	0.3784
13	11	1.0614	0.1637	0.5429
15	11	0.8848	0.1069	0.3547
16	7	1.0323	0.2414	0.6388
17	6	0.6609	0.2150	0.4808
18	9	0.3106	0.1261	0.3782
19	6	0.6032	0.1648	0.4038
20	8	0.5295	0.0778	0.2334
21	11	1.0217	0.1711	0.5675

Table 3.17. Transformed data for Peru F_2 generation female *N. prolixus*

Family number	n	\bar{x}	S.E.	S.D.
1	8	0.2634	0.0376	0.1064
2	4	0.8919	0.2259	0.4517
3	6	0.7564	0.1391	0.3407
4	8	0.4921	0.1510	0.4270
6	8	0.8917	0.0560	0.1583
7	5	0.9426	0.2708	0.6054
8	7	0.6460	0.1458	0.3856
9	6	0.3465	0.1360	0.3331
10	9	0.6724	0.1562	0.4686
11	7	0.8205	0.1387	0.3669
12	4	0.6276	0.1468	0.2936
13	8	0.7773	0.1779	0.5033
15	9	0.2928	0.1006	0.3019
16	7	0.9775	0.2384	0.6306
17	12	0.4998	0.1338	0.4633
18	7	0.5649	0.1707	0.4516
19	10	0.6808	0.1594	0.5042
20	7	0.4959	0.2053	0.5432
21	9	0.6748	0.1083	0.3249

Table 3.18. Transformed data for Peru F_2 families (male and female)
of R. prolixus

Family number	n	\bar{x}	S.E.	S.D.	Parental family number
1	18	0.5014	0.0916	0.3084	43
2	8	0.8013	0.1681	0.4756	43
3	15	0.7576	0.1179	0.4567	43
4	15	0.7113	0.1276	0.4942	63
6	15	0.8591	0.1012	0.3918	55
7	13	0.8275	0.1436	0.5176	55
8	12	0.8583	0.1436	0.4977	55
9	19	0.5621	0.0737	0.3214	56
10	19	0.7299	0.0776	0.3383	56
11	20	1.0339	0.0789	0.3530	5
12	13	0.7111	0.0863	0.3112	5
13	19	0.9418	0.1122	0.3855	5
15	20	0.6184	0.0990	0.4426	5
16	14	1.0049	0.1632	0.6105	61
17	17	0.5472	0.1114	0.4595	73
18	16	0.4218	0.1045	0.4181	73
19	16	0.6517	0.1111	0.4565	42
20	16	0.5148	0.0960	0.3819	48
21	20	0.8656	0.1108	0.4953	55

Parental family number - F_1 family from which parents of F_2 families were selected.

Table 2.3. Transformed data for strain 7 parental generation

S. maritima

Sex	n	\bar{x}	S.E.	S.D.
♂	81	0.8314	0.0081	0.6262
♀	36	1.0000	0.0071	0.6000
♂ + ♀	77	0.9560	0.0079	0.5934

Table 3.20. Transformed data for Strain 7 F_1 generation maleP. promelas

Family number	n	\bar{x}	S.E.	S.D.
50	20	1.3559	0.1849	0.8270
69	11	1.4781	0.2005	0.6651
93	14	1.1160	0.2281	0.8533
73	12	0.5382	0.1561	0.5407
94	21	1.4630	0.1035	0.4745
58	17	1.5263	0.1750	0.7214
46	16	1.2555	0.1748	0.6991
55	4	1.5588	0.3173	0.6347
70	7	1.4313	0.3007	0.7954
76	3	0.9830	0.6112	1.0587
66	13	1.4672	0.1441	0.5197

Table 3.21- Transformed data for Strain 7 F₁ generation femaleS. pomonae

Family number	n	\bar{Q}	\bar{P}_B	\bar{P}_W
40	16	0.9944	0.1666	0.6665
69	11	0.6919	0.1617	0.5363
93	8	0.9106	0.2249	0.6362
73	8	0.0376	0.0376	0.1064
94	15	1.0914	0.1384	0.5361
58	9	1.5609	0.0571	0.1713
46	10	1.1226	0.1562	0.6426
55	6	1.2872	0.3209	0.7860
70	11	1.8006	0.2063	0.7990
76	8	0.8255	0.1722	0.4870
66	8	1.1404	0.1832	0.5812

Table 3.2. Transformed data for Strain 7 *W.* families (male and female) of *E. proulxii*.

Family number	n	\bar{z}	$S_{\bar{z}}$	$S_{\bar{z}^2}$
57	36	1.1957	0.1100	0.1210
69	22	1.0850	0.1522	0.7138
93	22	1.0413	0.1646	0.7719
73	20	0.9379	0.1088	0.4866
94	36	1.3500	0.0853	0.5118
58	26	1.5414	0.1148	0.5856
46	14	1.1851	0.1154	0.6729
55	10	1.3958	0.2230	0.7051
70	22	0.7961	0.1911	0.8965
76	11	0.8685	0.1896	0.6290
66	21	1.3427	0.1160	0.5315

Table 3.23. Transformed data for Strain 7 F_2 generation male
B. prolixus

Family number	n	\bar{x}	S.E.	S.D.	Parental family number
1	7	1.1440	0.2387	0.6315	50
2	7	0.6468	0.1096	0.4486	50
3	13	1.5463	0.1483	0.5347	58
4	8	1.5444	0.2875	0.6132	58
5	15	1.2261	0.1356	0.5251	94
6	5	1.2393	0.1269	0.2839	94
7	10	1.3600	0.1604	0.5072	46 x 70
8	9	1.7106	0.2282	0.6847	70 x 46
9	12	1.3895	0.1328	0.4601	93
10	13	0.9369	0.1582	0.5705	93
11	10	0.9527	0.2517	0.7960	70
12	14	1.4922	0.0970	0.3629	73
13	10	0.8785	0.1915	0.6056	73
14	6	0.4762	0.2493	0.6106	73
15	12	1.1378	0.1839	0.6371	69
16	6	1.3303	0.1644	0.4026	76

Table 3.24. Transformed data for Strain 7 F₂ generation femaleS. prolisus

Family number	n	\bar{x}	S.E.	S.D.	Parental family number
1	10	0.6503	0.1672	0.5286	50
2	8	0.9028	0.0965	0.2729	50
3	13	1.3250	0.1914	0.6903	58
4	13	1.1154	0.1462	0.5270	58
5	6	0.9140	0.2380	0.5829	94
6	8	1.0489	0.1735	0.4908	94
7	7	0.9758	0.1378	0.3646	46 x 70
8	8	1.1017	0.1684	0.4764	70 x 46
9	8	0.6904	0.1751	0.4953	93
10	4	0.5909	0.2291	0.4581	93
11	9	0.5203	0.2904	0.7511	70
12	14	0.6913	0.0870	0.3255	73
13	11	0.5619	0.1818	0.6030	73
14	5	0.0602	0.0602	0.1346	73
15	8	0.9375	0.1864	0.5272	69
16	4	0.6694	0.2268	0.4535	76

Table 3.24. Transformed data for Strain 7 F_2 generation femaleH. prolixus

Family number	n	\bar{x}	S.E.	S.D.	Parental family number
1	10	0.6583	0.1672	0.5286	50
2	8	0.9028	0.0965	0.2729	50
3	13	1.3250	0.1914	0.6903	58
4	13	1.1154	0.1662	0.5270	58
5	6	0.9140	0.2380	0.5829	94
6	8	1.0489	0.1735	0.4908	94
7	7	0.9758	0.1378	0.3646	46 x 70
8	8	1.1017	0.1684	0.4764	70 x 46
9	8	0.6904	0.1751	0.4953	93
10	4	0.5909	0.2291	0.4581	93
11	9	0.5203	0.2904	0.7511	70
12	14	0.6913	0.0870	0.3255	73
13	11	0.5619	0.1818	0.6030	73
14	5	0.0602	0.0602	0.1346	73
15	8	0.9375	0.1864	0.5272	69
16	4	0.6694	0.2268	0.4535	76

Table 3.7. Transformed data: Not Omeas T_1 , families scale and female of B. proluxa

Family	n	\bar{x}	S.E.	S.E.	Parental family
1	17	0.8583	0.1470	0.6062	50
2	15	0.7833	0.0969	0.3755	51
3	26	1.4357	0.1000	0.6154	58
4	21	1.2789	0.1454	0.6661	58
5	21	1.1369	0.1193	0.5466	94
6	13	1.1221	0.1166	0.4203	94
7	17	1.2018	0.1170	0.4823	46 x 70
8	17	1.4240	0.1594	0.6571	70 x 46
9	20	1.1099	0.1227	0.5880	93
10	17	0.8555	0.1342	0.5534	93
11	19	0.7478	0.1802	0.7853	70
12	28	1.0918	0.1001	0.5299	73
13	21	0.7127	0.1333	0.6108	73
14	11	0.2871	0.1480	0.4908	73
15	20	1.0577	0.1318	0.5895	69
16	10	1.0659	0.1658	0.5245	76

Table 3.26. Transformed data for Strain 7 F_3 male M. proliger, based on the mean of two observations for each bug

Family number	n	\bar{X}	S.E.	S.D.	Selection lines	Parental family number
1	12	0.3274	0.0905	0.3133	1	10
2	16	0.6970	0.0689	0.2757	2	10
3	20	0.6999	0.0806	0.3603	3	11
4	8	1.1599	0.3114	0.6228	4	11
5	10	1.1603	0.0903	0.3833	5	10
6	8	0.9796	0.2933	0.6558	6	10
7	12	0.6849	0.1792	0.4276	7	10
10	11	0.6827	0.2045	0.4581	8	10
11	15	0.5841	0.1886	0.5988	9	10
12	14	1.4104	0.0993	0.4095	10	4
13	3	1.6855	0.8888	0.0449	11	5
14	20	1.5172	0.1885	0.2756	12	5
15	12	1.8025	0.1587	0.5428	13	5
16	10	1.6906	0.1888	0.4090	14	8
17	10	1.3956	0.1131	0.4483	15	8
18	16	1.870	0.2081	0.4149	16	8
19	8	1.2918	0.2637	0.6459	17	8

Key to selection lines for Strain 7 families:-

- 1 = selection for low score
- 2 = reverse selection from low-scoring families
- 3 = reverse selection from high-scoring families
- 4 = selection for high score

Table 3.27. Transformed data for Strain 71, female *A. pallipes*, based on the mean of two observations for each bug

Family number	N	T	1st	2nd	Selection line	Parental family number
1	11	0.3662	0.1131	0.1752	2	10
2	18	0.4097	0.0834	0.3536	3	6
3	11	0.4466	0.0850	0.2820	4	11
4	9	0.7438	0.1382	0.4146	5	10
5	19	0.6814	0.1083	0.4721	6	11
6	8	0.9904	0.1499	0.4241	7	10
7	6	0.7326	0.0718	0.1758	8	10
10	13	0.3644	0.1378	0.4967	9	10
11	15	0.4347	0.1508	0.5841	10	10
12	18	1.2190	0.1366	0.5795	11	9
13	5	1.3052	0.2360	0.5278	12	9
14	9	1.0102	0.1432	0.4295	13	9
15	12	1.7118	0.1455	0.5041	14	9
16	12	0.9960	0.1659	0.5746	15	8
17	8	1.0322	0.0704	0.1991	16	8
18	8	1.2303	0.1011	0.5405	17	8
19	4	1.4082	0.2134	0.4287	18	9

Table 7.28 Transformed data for Strain 7 F_3 families (male and female) of R. prolixus, based on the mean of two observations for each bug

Family number	n	\bar{X}	$S.E._x$	$S.E._y$	Selection line	Parental family number
1	48	0.3459	0.0702	0.3368	1	10
2	35	0.5449	0.0595	0.3467	1	10
3	41	0.6100	0.0632	0.3520	1	10
4	18	0.8719	0.1391	0.5015	1	10
5	37	0.8881	0.0805	0.4896	1	10
6	17	0.9862	0.1382	0.4983	1	10
7	18	0.7008	0.0880	0.3732	1	10
8	24	0.5103	0.1216	0.5959	1	10
9	30	0.5095	0.1070	0.5862	1	10
10	28	1.3119	0.0856	0.5062	1	10
11	11	1.4478	0.1575	0.4455	1	10
12	13	1.3789	0.0778	0.4467	1	10
13	24	1.2379	0.1188	0.4488	1	10
14	27	1.3763	0.1138	0.5915	1	10
15	28	1.2745	0.0851	0.4168	1	10
16	28	1.3948	0.0947	0.4640	1	10
17	31	1.3383	0.1721	0.5441	1	10

Table 1.2. Analysis of variance of male scores of parental generations of K. trilineus infected with T. parvi of 2 strains, Strain 7 and Peru.

Item	df	MS	VR	P
Between strains	1	2.5362	0.6433	0.01-0.02**
Within strains	81	0.2630		

df = degrees of freedom

MS = Mean square

VR = Variance ratio

Table 1.3. Analysis of variance of female scores of parental generations of K. trilineus infected with T. parvi of 2 strains, Strain 7 and Peru.

Item	df	MS	VR	P
Between strains	1	6.7611	26.3283	<0.001***
Within strains	74	0.2568		

Table 1.4. Analysis of variance of male and female scores of parental generations of K. trilineus infected with T. parvi of 2 strains, Strain 7 and Peru.

Item	df	MS	VR	P
Between strains	1	8.5730	32.6839	<0.001***
Within strains	156	0.2623		

These 3 analyses show that in the parental generation there were highly significant differences in intensity of infection between bugs infected with the 2 difference strains of trypanosome, when compared to variation between bugs within strains and that these differences were common to both sexes.

A similar analysis was carried out on the data from F_1 bugs infected with different strains of trypanosome (Table 3.32).

Table 3.32. Analysis of variance of male and female parents of F_1 generations of H. prolixa infected with T. cruzi of 2 strains, strain 7 and 7000.

Item	df	SS	MS	P
Between strains	1	20.4691	10.1679	<0.001**
Between bug families (within strains)	21	1.6811	5.3267	<0.001***
Between individual bugs (within families)	241	0.3156		

This analysis shows that there were highly significant differences in intensity of infection between F_1 bugs infected with the 2 strains of T. cruzi, even when this variation was tested against the considerable differences between F_1 families. Comparison of further generations was not made as such analyses would not be strictly valid because of the different amounts of selection pressure which were applied to subsequent generations of the 2 groups of bugs. However, it is clear that differences between T. cruzi strains were very large and these are reflected in the

mean scores for the parental generations, shown in Tables 3.12 and 3.19: Peru mean = 0.49, S.E. = 0.03; Strain 7 mean = 0.96, S.E. = 0.07. It may be concluded that bugs infected with Strain 7 T. cruzi excreted much greater numbers of trypanosomes (bearing in mind that these scores have been transformed logarithmically) than those infected with the Peru strain. This result is even more remarkable when it is recalled that bugs feeding on mice infected with Peru strain were ingesting more parasites than the bugs feeding on mice infected with Strain 7, the parasitaemia of the Strain 7 mice being <1 trypanosome per field of tail blood at the time of feeding, while the Peru mice had 1-5 trypanosomes per field. These differences suggest strongly that the intensity of infection of an infected bug is markedly influenced by the genotype of the trypanosome ingested. The significant difference demonstrated between bugs of the parental generations is important since these bugs were selected at random from the same colony, and prior to selection may be assumed to have been genetically homogeneous, suggesting that differences between parental groups are a consequence of genotypic differences between strains of trypanosome. However, the highly significant differences between F_1 families within T. cruzi strains when tested against the variation between individual bugs within families further suggests that the bug genotype (determined by family differences), as well as trypanosome genotype, is involved in determining the intensity of infection in H. prolixus.

Differences between sexes and families

The variation in intensity of infection between sexes of H. prolixus was analysed for both Strain 7 and Peru infections in bugs of all generations.

1. Parental generation.

Differences between male and female scores for bugs of the parental generation, were compared by analysis of variance, the results of which are shown in Tables 3.33 and 3.34.

Table 3.33. Analysis of variance of male and female scores for parental generation *H. prolixus* infected with *T. cruzi* Peru strain.

Item	df	MS	VR	P
Between sexes	1	0.1488	—	N.S.
Within sexes	78	0.0708		

Table 3.34. Analysis of variance of male and female scores for parental generation *H. prolixus* infected with *T. cruzi* Strain 7.

Item	df	MS	VR	P
Between sexes	1	1.3610	4.0183	0.05-0.01*
Within sexes	78	0.3387		

These results show that in the parental generation there were no significant differences between male and female scores for bugs infected with Peru strain while there were significant differences between sexes for bugs infected with Strain 7 *T. cruzi* when compared to the variation between individuals within each sex.

2. F₁ generation

(a) Peru strain. The mean male score in F₁ generation for male bugs infected with T. cruzi Peru strain was greater than the mean female score (\bar{x} males = 0.83, S.E. \bar{x} = 0.08; \bar{x} females = 0.68, S.E. \bar{x} = 0.21).

These differences in mean score were examined using analysis of variance, the results of which are shown in Table 3.35.

Table 3.35. Analysis of variance of male and female scores for F₁ generation T. cruzi infected with T. cruzi Peru strain.

Item	df	SS	MS	P
Between sexes	1	9.7453	9.7453	0.05-0.1 ⁰
Increment due to families	29	1.3644	0.4705	0.1***
Residual	283	0.1937	0.0007	

The results show that there were slightly significant differences between sexes when compared to the variation between families. The increment of variation due to families was highly significant compared to the variation between individuals within one family. This variation between one-family was partitioned and a further analysis carried out, the results of which are shown in Table 3.36.

Table 3.36. Analysis of variance of male and female scores for χ_1 generation K₁ broilers infected with T₁ Peru strain.

Item	df	MS	VK	P
Between sexes	1	0.7453	Between	0.01-0.05*
Between families	11	0.9697	5.0052	<0.001***
Sex X families	11	0.2120	1.0944	N.S.
Residual	281	0.1937		

This analysis shows that variation between sexes was significant when tested against the error item. The variation between families was highly significant when tested against the error and there was no significant interaction between sex and family, indicating that the difference between sexes was constant for all the 12 families infected with Peru strain T₁ Peru.

(b) Strain 7. The mean score of χ_1 males was greater than that of females infected with Strain 7 (\bar{x} males = 1.31, S.E. \bar{x} = .44; \bar{x} females = 0.94, S.E. \bar{x} = 0.31).

This difference was again examined by analysis of variance, the results of which are shown in Tables 3.37 and 3.38.

Table 3.37. Analysis of variance of male and female scores for χ_1 generation K₁ broilers infected with T₁ Peru Strain 7.

Item	df	MS	VK	P
Between sexes	1	6.3751	4.7612	0.01-0.05*
Increment due to families	20	1.3390	3.1714	<0.001***
Residual	238	0.4222		

Table 3.38. Analysis of variance of male and female scores for P_1 generation K. pulikow infected with T. cruzi Strain 7, partitioning the variation due to families.

Item	df	SS	VK	P
Between sexes	1	6.3751	15.0994	<0.001***
Between families	10	2.1975	5.2047	<0.001***
Sex X families	10	0.4340	1.0379	N.S.
Residual	238	0.4222		

The results in Table 3.37 show that the differences between sexes were significant when tested against the variation due to families, but when this increment due to families was partitioned (Table 3.38) the variation between sexes was highly significant when tested against the residual variation. Table 3.38 also shows that there was no significant interaction between sex and family when this item was tested against the error. Indicating that differences between sexes were consistent for all families of this generation. As with the T. cruzi Peru strain infected bugs, differences between families in Stomoxys infection were highly significant when tested against the error for P_1 bugs infected with T. cruzi Strain 7.

3. P_2 generation

(a) Peru strain. The mean score of P_2 males infected with Peru strain T. cruzi was greater than the mean score for females infected with the same strain (\bar{x} males = 0.81, S.E. \bar{x} = 0.19; \bar{x} females = 0.63, S.E. \bar{x} = 0.14). These differences were again examined by analysis of variance, the results of which are shown in Tables 3.39 and 3.40.

Table 3.3¹. Analysis of variance of male and female scores for F_2 generation *M. prolixus* infected with *T. cruzi*, Peru strain.

Item	df	MS	VR	P
Between sexes	1	1.7408	4.6423	0.05-0.1*
Increment due to families	36	0.0750	2.0470	0.00000000**
Residual	267	0.1832		

Table 3.4¹. Analysis of variance of male and female scores for F_2 generation *M. prolixus* infected with *T. cruzi*, Peru strain, partitioning the variation due to families.

Item	df	MS	VR	P
Between sexes	1	1.7408	4.6423	0.05-0.1**
Between families	18	0.4870	0.0750	0.00000000***
Sex x families	18	0.2419	1.3203	N.S.
Residual	267	0.1832		

The results in Table 3.39 show that the differences between sexes were significant when tested against the variation due to families and by partitioning this last item (Table 3.40) the differences between sexes were shown to be highly significant when tested against the error variation. The variation between F_2 families was highly significant when tested against the error variation (Table 3.40) but there was no

significant interaction between sex of bug and family, again showing that sex differences were constant over families.

(b) Strain 7. The mean scores for the two sexes of *P.* bugs again differed with this strain of trypanosome; the males having the higher mean score as in previous generations (\bar{X} males = 1.77, S.E. = 0.04; \bar{X} females = 0.74, S.E. = 0.03), and this difference was tested by analysis of variance, the results of which are shown in Tables 3.41 and 3.42.

Table 3.41. Analysis of variance of male and female scores for *P.* generation, *he. trilineus* infected with *S. traw* strain 7.

Item	df	MS	VR	P
Between sexes	1	9.8183	11.3665	<0.001***
Increment due to families	30	0.8638	2.8770	<0.001***
Residual	261	0.3002		

The difference between sexes was highly significant even when tested against the large increment of variation due to families, but this increment was again partitioned to test for any sex X family interaction.

Table 3.4. Analysis of variance of male and female scores for F_2 generation H_2 population infected with T. cruzi Strain 7, partitioning the variation due to families.

Item	df	MC	VR	P
Between sexes	1	9.8183	32.7016	<0.001***
Between families	18	1.4278	8.4891	<0.001***
Sex X families	18	3.2775	10.9247	N.S.
Residual	361	0.3002		

Differences between F_2 families were highly significant in both analyses but the sex X families interaction was not significant, demonstrating once more that differences between sexes were constant for families of different pedigree. The highly significant differences between families which are a constant feature of the analysis of intensity of infection shown above strongly suggest that bug genotype is involved in regulating trypanosome growth in the host. These family differences were demonstrated for both trypanosome strains used and were independent of the differences, since no significant sex X family interaction terms were found on analysis of data. Differences in variance between families must reflect the similarity of within-family scores which can only have resulted from the genetic resemblance of siblings.

4. F_3 generation

Only bugs infected with Strain 7 T. cruzi were bred to the F_3 generation. An analysis of variance of male and female scores was

carried out, the results of which are given in Table 3.43.

Table 3.44. Analysis of variance of male and female scores for F_1 generation *A. proxima* of 2 selection lines from 17 families infected with *T. virus* Strain 7, each score based on the mean of 2 observations.

Item	df	SS	MS	F
1. Between sexes	1	10.9109	10.9109	<0.001***
2. Sex X lines	1	8.091	8.091	<0.001***
3. Sex X families (within lines)	15	0.2732	-	N.S.
4. Residual	374	0.3613	-	-
1 + 4. Pooled	384	0.3579	-	-

The results show that differences between sexes were highly significant when tested against items 3 and 4 pooled. There was no significant sex X family interaction, even more indicating that the significant differences between sexes were constant for different families. The sex X lines interaction (item 2) was highly significant when tested against the error variation indicating that differences between sexes was affected by the direction of selection. A survey of the mean values for the F_1 generation shows that while the mean male score ($\bar{X} = 1.11$) was greater than the female mean ($\bar{X} = 0.78$) for all families, the difference between sexes was greater in families

numbered 12 - 19 of the high selection lines (\bar{x} males - \bar{x} females = 0.13) than between families numbered 1 - 11 of the low selection lines (\bar{x} males - \bar{x} females = 0.04). It may be that this difference is an effect of scaling, and that the logarithmic transformation did not completely normalize the data or, alternatively, it could be that control of susceptibility is determined by a sex-linked gene or genes.

Since throughout the experiment the results have shown that male scores for trypanosomes in their faeces were greater than female scores, several possible causes, other than bug or trypanosome genotype were tested. First, the explanation for higher levels of male infections may have been simply that fifth instar males (the stage at which the bugs, male and female, were given their infecting meal throughout these experiments) were taking larger infecting blood-meals than female larvae. This possibility was investigated by weighing 2 groups of fifth instar larvae of different sex before and after a blood-meal, and comparing the weights of the blood-meals. The results of this experiment are shown in Table 3.44.

Table 3.44. Analysis of blood-meal weights for 20 male and 20 female fifth instar K. bollinus fed on mice. Weights in grams.

		Mean weight of unfed bugs	Mean weight of fed bugs	Mean weight of blood-meal
Males	\bar{x}	0.1081	0.1815	0.0734
	S.E. \bar{x}	0.0009	0.0041	0.0033
Females	\bar{x}	0.0257	0.1089	0.0832
	S.E. \bar{x}	0.0011	0.0067	0.0033
t		1.1959	1.8398	1.0725
P		N.S.	N.S.	N.S.

\bar{x} = Mean; S.E. = Standard error

The results of the t-tests show that there was no significant difference between the amount of blood taken by male and female fifth instar larvae, and it may be inferred that experimental bugs of different sex took approximately the same size infecting blood-meals.

Secondly the relationship between sex and trypanosome excreta was further investigated by comparing the rate of defaecation of male and female adult E. tritritus. For this comparison, male and female adults (which, like the experimentally infected bugs, had not had a previous meal as adults) were individually fed to repletion and then each allowed to excrete for one hour, when the weight of excreta collected in the tubes was calculated by difference in weight. The results of this experiment are shown in Table 3.45.

Table 3.45. Analysis of blood-meal and excreta weights in 18 male and 18 female E. tritritus adults. Weights in grams.

	Mean weight of unfed bugs	Mean weight of fed bugs	Mean weight of blood-meal	Mean weight of excreta after 1 hr.	% Blood-meal excreted in 1 hr.
Males \bar{x}	0.0509	0.1411	0.0900	0.0906	26.35
S.E. \bar{x}	0.0019	0.0046	0.0031	0.0035	
Females \bar{x}	0.0681	0.2031	0.1354	0.0054	40.01
S.E. \bar{x}	0.0033	0.0075	0.0046	0.0054	
t	3.8252	6.7063	7.9208	3.3056	
P	<0.001***	<0.001***	<0.001***	0.01-0.001**	
χ^2					16.2082
P					<0.001***

\bar{x} = Mean

S.E. \bar{x} = standard error of mean

Weight excreta = weight of excreta collected in 1 hr. post feeding

$$\% \text{ blood-meal excreted} = \frac{\text{weight excreta}}{\text{weight blood-meal}} \times 100$$

The results of the t-tests between sexes show that adult females took significantly larger blood-meals than males and also excreted significantly larger amounts of faeces than males within one hour after feeding. The χ^2 test for proportions showed that females also excreted a greater proportion of their blood-meal within one hour of feeding than did the adult males. Given these results, it would be expected that, on an infecting blood-meal, fifth instar males and females would ingest approximately the same numbers of trypanosomes and, when subsequently given a clean feed, the females would excrete a greater number of trypanosomes within a limited period. However, the results of the present work have demonstrated the contrary; males excreted significantly more trypanosomes than adult female *P. tritici* throughout the experiment. It may be concluded that differences in intensity of infection between sexes are independent of variations in blood-meal size (and thereby numbers of trypanosomes ingested) and of differences in rates of excretion, since females took larger blood-meals and yet excreted significantly fewer trypanosomes. These sex differences appear to be independent of environmental variables and must be attributed to genetically determined differences between the two sexes of *P. tritici*.

Differences between selection lines

In analysing the effects of selection for intensity of infection in successive generations of *P. tritici*, only data for bugs infected with Strain 7 (Table 1, 12) = 1,284 were used and these bugs strain infected bugs were only bred to the F_2 generation and could not provide sufficient information for a reasonable assessment of the changes brought about by selection. The bugs of the F_1 generation infected with Strain 7 were each scored twice by repeated sampling of

The results of the t-tests between sexes show that adult females took significantly larger blood-meals than males and also excreted significantly larger amounts of faeces than males within one hour after feeding. The χ^2 test for proportions showed that females also excreted a greater proportion of their blood-meal within one hour of feeding than did the adult males. Given these results, it would be expected that, if an infecting blood-meal, either from a male or female, would ingest approximately the same numbers of trypanosomes and, when subsequently given a clean feed, the females would excrete a greater number of trypanosomes within a limited period. However, the results of the present work have demonstrated the contrary: males excreted significantly more trypanosomes than adult female H. prolixus throughout the experiment. It may be concluded that differences in intensity of infection between sexes are independent of variations in blood-meal size (and thereby numbers of trypanosomes ingested) and of differences in rates of excretion. Since females took larger blood-meals and yet excreted significantly fewer trypanosomes. These sex differences appear to be independent of environmental variables and must be attributed to genetically determined differences between the two sexes of H. prolixus.

Differences between selection lines

In analysing the effects of selection for intensity of infection in successive generations of H. prolixus, very data for long collected with T. tritici Strain 7 (values 1.19 - 1.28) have been used since these strain infected bugs were only bred to the 7th generation and could not provide sufficient information for a reasonable assessment of the changes brought about by selection. The bugs of the 7th generation infected with Strain 7 were each scored twice by repeated sampling of

their diluted faeces and analyses for this generation have been based on the mean of these 2 observations. The effects of selection pressure on mean family scores over successive generations of bugs infected with T. cruzi Strain 7 are illustrated in Figs. 3.3 - 3.8. In these diagrams the mean family scores have been plotted for each generation of selection and assortative sib-mating, the pedigree of each family being traced by the lines connecting the means. The numbers beside the P_3 means correspond to the P_3 family code numbers in Tables 3.26 - 3.28.

Figs. 3.3 and 3.4 show the changes in mean family score for both sexes, Fig. 3.3 showing results of selection for low scores and Fig. 3.4 showing results of selection for bugs with high intensity of infection. Comparison of these 2 figures shows that family means did change in response to selection pressure for both high and low scores. However, since significant differences between sexes have already been demonstrated, it is more meaningful to plot response to selection of family means with sexes treated separately and these changes are illustrated in Figs. 3.5 - 3.8. These diagrams show that both sexes of Strain 7 infected bugs responded to selection pressures for high and low scores but they also reflect the differences between sexes demonstrated statistically above. For example, a comparison of P_3 family 12 mean in Fig. 3.7 with the mean value for females of the same family in Fig. 3.8 shows that the males responded to assortative selection pressure, but the females of the same family did not. However, a closer inspection of the results in Tables 3.26 and 3.27 shows that the male score value was based on scores from 6 males while the female value was based on only 4 females. In fact, family 19 was one of the smallest P_3 families and therefore too much emphasis cannot be placed on this particular family, but it serves to emphasise that the mean values shown in

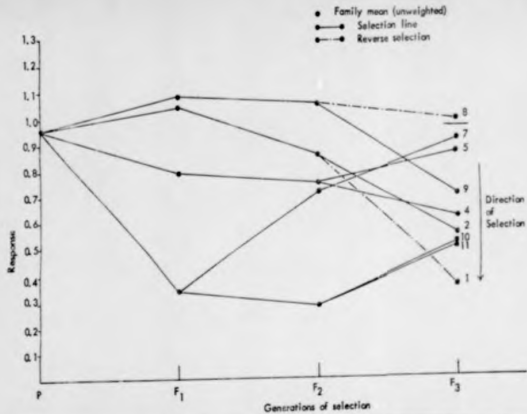


Figure 3.3. Changes in family means in response to selection for low levels of infection in *S. prolixus* infected with *T. cruzi* Strain 7.

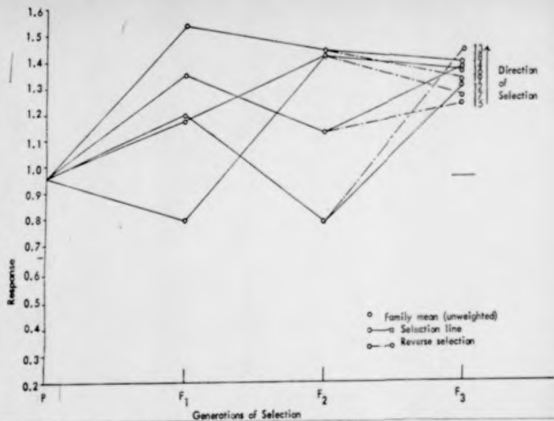


Figure 3.6. Changes in family means in response to selection for high levels of intensity of infection in *R. prolixus* infected with *T. cruzi*.

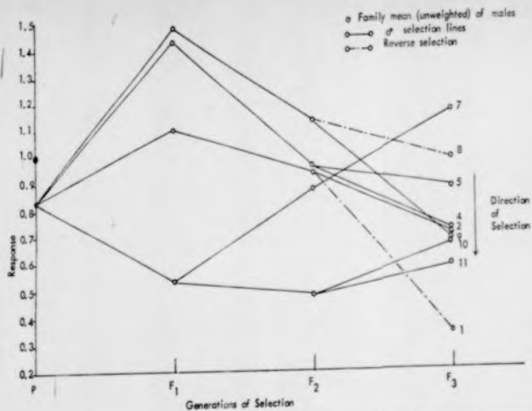


Figure 3.5. Changes in mean male scores in response to selection for low levels of intensity of infection in *R. prolixus* infected with *T. cruzi* Strain 7.

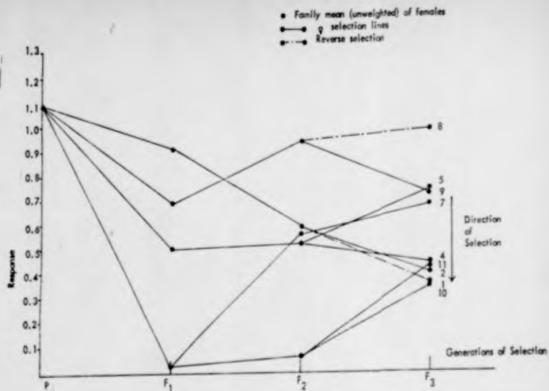


Figure 3.6. Changes in mean female scores in response to selection for low levels of intensity of infection in B. prolixus infected with T. cruzi Strain 7.

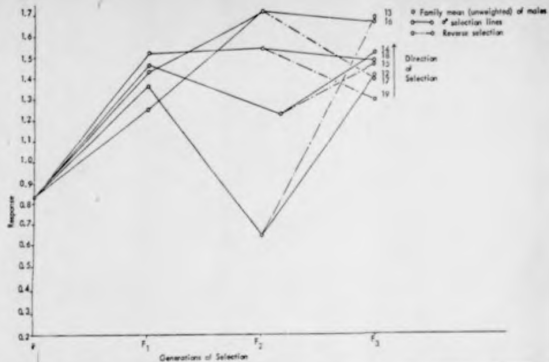


Figure 3.7. Changes in mean male scores in response to selection for high levels of intensity of infection in *B. prolixus* infected with *T. cruzi* Strain 7.

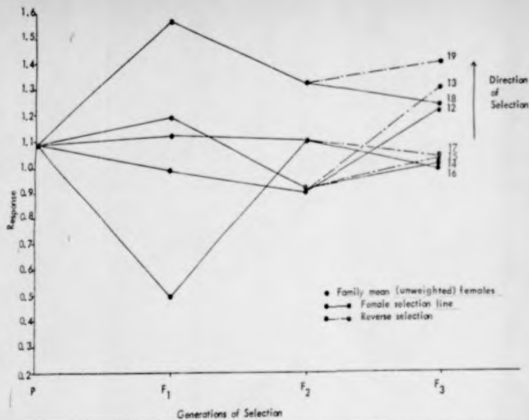


Figure 3.8. Changes in mean female scores in response to selection for high levels of intensity of infection in *E. prolixus* infected with *T. cruzi* Strain 7.

Figs. 3.1 - 3.6 do not convey a very meaningful impression of the response of family means to selection; firstly the means are based on differing numbers of individuals but are not weighted to compensate for this, and secondly, these diagrams give no impression of the degree of selection applied to each family.

However, differences between F_3 families infected with is. vir. Strain 7 have been quantified, by using analysis of variance, which gives a meaningful comparison because sum of squares are calculated in relation to the number of observations in each family. The differences between males of F_3 families selected for high and low intensity of infection were examined by an analysis of variance, the results of which are shown in Table 3.46.

Table 3.46. Analysis of variance of male scores of 11 F_3 generation families infected with is. vir. Strain 7 from 2 selection lines. Each score based on the mean of 2 observations.

Item	df	MS	Vk	P
Between selection lines	1	30.5385	58.0911	0.0000***
Between families (within lines)	15	2.3287	2.6031	0.01-0.001**
Between individuals (within families)	20	0.2034		

The results of this analysis show that there were highly significant differences in intensity of infection between males selected for high and low levels of intensity of infection when tested against the variation between families within selection lines. The variation between families

was also significant when tested against the variation between individuals within families.

A similar analysis of variance was carried out on females of the F_2 generation, the results of which are shown in Table 3.47.

Table 3.47. Analysis of variance of female scores of 17 F_2 families infected with 1.1001 strain 7 from 2 selection lines. Each score based on the mean of 4 observations.

Item	df	MS	VR	P
1. Between selection lines	1	15.2806	44.400	0.0001***
2. Between families (within lines)	15	3.3438	9.507	0.0001***
3. Between individuals (within families)	16	0.2157		
Between selection lines	1	15.2806	44.400	0.0001***
Pooled SS for 2 and 3	184	0.2262		

This analysis of female scores shows that there were highly significant differences in intensity of infection between bugs selected for high and low scores when compared to the variation between families within selection lines. The variation between families within lines was slightly significant when compared to the variation between individuals within families, and these 2 items were pooled to form further the differences between selection lines, and again the differences between

selection lines were shown to be highly significant. It may be concluded from these analyses that, as a result of selection by assortative sib-mating from a single parental population, 2 groups of families were produced in the F_3 generation, one of which contained bugs with high levels of intensity of infection, the other bugs with low levels, and that differences between these 2 groups were highly significant. Thus the differences between the 2 groups of F_3 families illustrated in Figs. 3.3 - 3.8 represent real differences in both sexes of bug infected with T. cruzi Strain 7. A further comparison was made between the F_3 family with the highest mean score from the families of the low selection lines (family 7; Fig. 3.3) - ignoring reverse selected families - and the family with the lowest mean score from families selected for high scores (family 12; Fig. 3.4). The results of this analysis are given in Table 3.4b.

Table 3.4b. Analysis of variance of F_3 families 7 and 12 from 2 selection lines for high and low score.

Item	df	MS	VK	P
Between families	1	2.8437	11.4776	0.01-0.001**
Within families	70	0.2477		

The results show that there was a highly significant difference between these 2 families when tested against the variation between individuals within families which is reflected in the means: family 7 $\bar{x} = 0.9144$, S.E. = 0.0088; family 12 $\bar{x} = 0.5419$, S.E. = 0.0088. $F_{1,70} = 11.48$, $P < 0.01$.

analysis provides further evidence of the efficacy of the two-way selection programme which produced 2 clearly separated groups of families by the F_3 generation.

Reverse selection.

Attempts were made to reverse the selection process from some F_2 families and these families are indicated by broken lines in Figs. 3.1 - 3.8. Owing to mortality, only 7 reverse selection matings were successful from the low selection lines (F_2 families 1 and 8), and comparisons between these families and the other 7 families of the same selection group have not been made since insufficient reverse selected control data was available for a meaningful analysis. However, from each F_2 family of the high selection group, 3 matings were established, firstly a mating selected for high score and secondly a mating reverse selected for low score. A fair comparison can be made of these 7 groups, each of 4 families, and an analysis of variance has provided the basis of this data, the results of which are shown in Tables 3.4a and 3.4b (see below) and discussed respectively.

Table 3.4a. Analysis of variance of male scores for F_2 families 12, 15, 16 and 18 of high selection lines and families correspondingly reverse selected for low scores, 13, 15, 17 and 19.

Item	df	MC	VR	P
Between lines	1	0.2154	1.4875	0.05
Between families (within lines)	6	0.1598	—	N.S.
Between individuals (within families)	101	0.1204	—	—

Table 3.11. Analysis of variance of female scores for F_3 families 13, 14, 16 and 18 of high selection lines and families correspondingly reverse selected for low score, 13, 15, 17 and 19.

Item	df	MS	Vk	F
Between lines	1	0.0	-	N.S.

These analyses show that there were no significant differences between bugs of either sex of the high selection lines and those families reverse selected for low score for general performance. However, it must be noted that these families were inbred as a result of self-mating to F_3 and therefore the reverse selected and control families would be genetically similar, which may account for the similarity of means in the 2 groups. But before any conclusive interpretation could be based on these results, further generations of reverse selection would have had to be carried out.

Having demonstrated in analyses in Tables 3.10, 3.11 and 3.12 that selection for high and low scoring bugs was successful by the F_3 generation, it is necessary, to gain an insight into these results, to relate the amount of selection pressure applied to the response of the selected families.

Measurement of response to selection

1. Definition of response to selection. The change in population mean brought about by selection, is called the response, but when more than one

generation of selection has been made, the measurement of the response is not simple. Inspection of Figs. 1.1 - 1.8 shows that the generation means of the families infected with Strain 7 did not progress in a regular fashion but fluctuated erratically. This is a phenomenon common to most selection programmes and Falconer (1967) attributes it to two causes: firstly, sampling variation which is determined by the number of individuals measured and, secondly, environmental variation. In the present experiment the sampling variation was large, variable numbers of bugs being raised in each family, and environmental variation may also have affected the results, for although temperature and humidity were closely regulated, other factors outlined in Fig. 3.1 may have affected the results, if only in a minor way. Where there is variation in generation means, Falconer (loc. cit.) states that the best measure of the average response per generation is obtained by fitting a regression line to the means, thereby assuming that the true response has been constant over the selection programme. The use of a two-way selection procedure in the present experiment allowed for a more accurate measurement of the response to selection to be made than would have been obtained from a comparison with an unselected control population, since environmental changes equally affected both selection lines. The response to selection measured from the divergence of the two regression lines will be about twice that of the lines separately, each line acting as a control for the other.

The response (R) to selection is defined as the difference in mean phenotypic value between the offspring of the selected parents and the whole of the parental generation. The measure of the selection

applied is the average superiority of selected parents and is called the selection differential (S) which is defined as the mean phenotypic value of selected parents expressed as a deviation from the population mean before selection (Falconer, loc. cit.). The regression of offspring score on parents is thus R/S and therefore $R = b_{op} S$ where b_{op} is the regression (b) of offspring on mid-parent value (op). Having carried out a selection programme, the equation of response given above can be used to estimate the heritability of a character from the results of the selection experiment. It can be shown that $R_{reg} = h^2 S$ (where h^2 is the heritability) thus $h^2 = R/S$, this ratio being defined as the realized heritability (Falconer, loc. cit.) and providing a measure of the effectiveness of selection. Realized heritability may be measured by plotting response against selection differential and, to obtain a measure of this differential that is relevant to families of different size, the selection differential must be weighted according to the numbers of progeny scored. The effective selection differentials, appropriately weighted, may then be carried over successive generations to give the total selection applied up to any given generation and this cumulated selection differential may then be plotted against the response. A regression line is then fitted to the points and the slope of this line measures the average value of R/S the realized heritability (Falconer, loc. cit.).

2. Estimates of heritability. To determine the realized heritability of susceptibility to infection from the present work, cumulated selection differentials and response statistics, as defined above, were calculated for families infected with *P. aureus* strain T₃, the results of which are shown in Tables 3.51 and 3.52 for families selected for low scores and high scores respectively (reverse selections were excluded). Weighted regression analyses were carried out using the R and S values given in these

Table 3.51. Cumulated selection differentials (C.S.), response (R) and weightings (WT) for families selected for low score, infected with T. cruzi strain 7.

Generation	Family number	WT	C.S.	R
F ₁	93	22	-0.9560	0.0853
F ₂	10	17	-1.7615	-0.1005
F ₃	2	24	-2.5670	-0.4111
F ₁	70	22	-0.5045	-0.1599
F ₂	11	19	-1.0790	-0.2082
F ₃	4	31	-2.5225	-0.3460
F ₃	6	13	-2.5225	-0.0841
F ₁	73	20	-0.3540	-0.6181
F ₂	13	21	-0.7080	-0.2433
F ₃	7	17	-1.0620	-0.0416
F ₁	69	22	-0.2103	0.1290
F ₂	15	20	-0.4206	0.1017
F ₃	8	18	-1.1266	-0.2552
F ₁	73	20	-0.3540	-0.6181
F ₂	14	11	-0.7780	-0.6689
F ₃	10	24	-1.0620	-0.4457
F ₃	11	30	-1.0620	-0.4465

WT = Number of individuals scored per family

From scores of parental generation = 3.0000

Table 3.25. Cumulated selection differentials (S.E.I.), response (R) and weightings (WT) for families selected for high score, infected with T. cruzi strain 7

Generation	Family (number)	WT	S.E.I.	R
P_1	50	36	-0.6550	0.2392
F_1	5	15	1.1469	-0.1677
F_2	12	15	2.7795	0.3558
P_2	94	35	-0.8055	0.3940
F_2	5	21	1.4942	0.1803
F_3	14	33	2.2196	0.4229
P_3	70	22	-0.5045	-0.1559
F_4	8	17	1.1757	0.4680
F_5	16	27	2.2501	0.4203
P_4	58	26	-0.8002	-0.1964
F_6	3	26	0.8454	0.4797
F_7	18	24	2.2366	0.4388

WT = Number of individuals scored per family

Mean score of parental generation = 0.0000

Tables and the results have been expressed graphically in Figs. 3.4 and 3.10, in which the best fitting lines have been drawn in accordance with the regression analyses. These weighted regression analyses were computed according to the following equations:-

$$SS \text{ (sum of squares) of } x = \sum (wx - \bar{w}\bar{x})^2$$

$$SSy = \sum (wy - \bar{w}\bar{y})^2$$

$$\text{(Sum of products) } SP \text{ (xy) } = \sum (wx - \bar{w}\bar{x})(wy - \bar{w}\bar{y})$$

$$\text{Regression } SS = SP \text{ (xy)}^2 / SSx$$

$$\text{Total } SS = SSy$$

Where w = weight

$$x = \text{Time}$$

$$y = k$$

$$n = \text{no. pairs of observations}$$

The regression lines were tested by analysis of variance, the results of which are shown in Tables 3.52 and 3.54.

Table 3.52. Analysis of variance of regression lines for animals selected for low levels of intensity of infection with T. citri Strain 7.

Item	df	MS	VR	P
Regression	1	44.8770	15.1085	0.001 ***
Residual	369	1.3992		

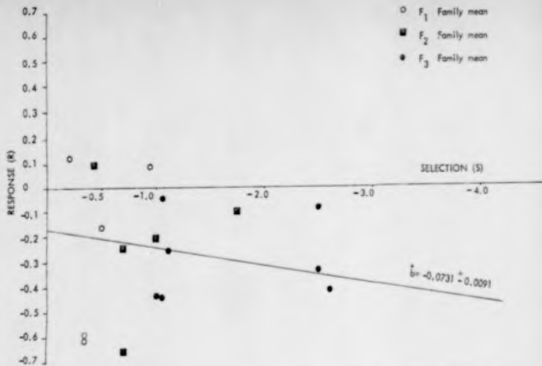


Figure 3.9. Response (R) to selection (S) for low levels of intensity of infection in *B. prolixus* infected with *T. cruzi* Strain 7.

R = Mean family score minus mean score of parental generation.

S = Cumulated selection differential (weighted).

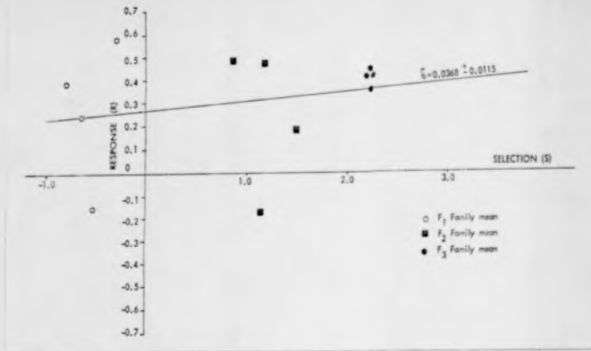


Figure 3.10. Response (R) to selection (S) for high levels of intensity of infection in B. pulex infected with T. cruzi Strain 7.

Table 3.b.i. Analysis of variance of regression items for families selected for high levels of intensity of infection with T. cruzi Strain 7.

Item	df	SS	VR	P
Regression	1	427.44	147.7735	.1***
Residual	316	2,892.9		

The results of these analyses show that the regression clearly accounts for practically all of the variance in both analyses, the remainder (non-linear) item being insignificant in both cases. These analyses show conclusively that response changed systematically with the selective pressure applied in both upward and downward directions.

The regression line for downward selection (Fig. 3.a) had a slope (b) of -0.701 ± 0.04 and, since this regression was highly significant, it may be concluded that the realized heritability for bugs selected for reduced levels of intensity of infection was 0.49 ± 0.06 . The regression line for upward selection (Fig. 3.b) had a slope (b) of 0.666 ± 0.045 and, since this regression was also highly significant, it may be concluded that the realized heritability for *H. tritrix* selected for high levels of intensity of infection was 0.38 ± 0.07 . From these two estimates, it was concluded that the mean realized heritability for this character was 5.0%.

Reversion Selection:

Although an analysis of variance of mean F_3 family scores (Tables 3.4c and 3.5d) failed to show any differences for reversion selected families, that analysis did not allow for the amount of selection

Table 3.55. Cumulated selection differentials (C.F.), response (R) and weightings (WT) for families reverse selected from high scoring lines infected with T. cruzi Strain 7.

Generation	Family number	WT	C.S.	R
F ₁	50	36	-0.6550	0.2392
F ₂	2	15	1.1469	-0.1677
F ₃	13	8	0.8520	0.4918
F ₁	94	36	-0.8055	0.3940
F ₂	5	21	1.4942	0.1803
F ₃	15	24	0.8777	0.2819
F ₁	46	34	-0.1779	0.2291
F ₂	8	17	1.5647	0.4680
F ₃	17	24	0.8572	0.3185
F ₁	58	26	-0.3055	0.5845
F ₂	3	26	0.8454	0.4797
F ₃	19	10	0.6645	0.3823

pressure applied to the 4 groups of families. The calculated selection differentials and response values have been calculated for F_3 families 12, 13, 17 and 18, and are set out in Table 3.11 together with the appropriate weighting. The resulting weighted regression analysis showed a slope (b) of -0.1181 ± 0.0025 i.e. the regression lines were tested by analysis of variance, the results of which are shown in Table 3.16.

Table 3.16. Analysis of variance of regression items for families reverse selected from high lines.

Item	df	MS	VR	P
Regression	1	2.9495	-	N.S.
Residual	275	3.1875		

The results show that the regression item was not significant indicating that reverse selection from F_3 families did not produce the systematic increase in family mean demonstrated above for F_3 families 12, 13, 14 and 18 and that the relationship of families 12, 13, 17 and 18 in response to selection was not linear. This suggests that reverse selection was effective in disrupting the linear response to selection for increasing levels of intensity of selection.

4. Summary of response to selection. The results illustrated in Figs. 3.9 and 3.11 were here put together for comparative purposes in Fig. 3.12 which illustrates the comparability of response to selection in two directions achieved in the present experiment: selection for low

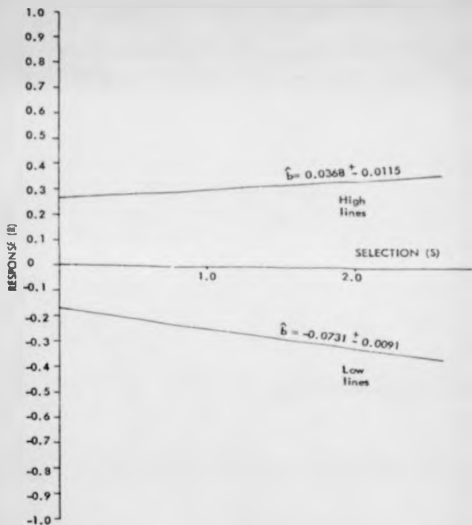


Figure 3.11. Asymmetry of response (R) to selection (S) for high and low levels of intensity of infection in *P. gouldii* infected with *T. cruzi* strain 7.

levels of intensity of infection was approximately twice as effective as selection for high levels. This asymmetry of response to selection has been found by different workers in selection experiments with other organisms and its possible causes will be discussed further.

DISCUSSION

The purpose of these genetic investigations on tritomine bugs was to determine what genetic mechanisms might be controlling their susceptibility to infection with the trypanosome, T. cruzi, for which they are natural vectors to man and other vertebrates in South (and parts of Central) America. This study was undertaken only with E. prolixus (infected with 2 strains of T. cruzi) as this species had the shortest generation time (approximately 8-6 months) of the species kept at the U.S.N. and T.R. Moreover, this species was available in colony in large numbers and, of importance, some earlier work by others had touched on the possibility of genetic inheritance of susceptibility in this species, an important vector in Venezuela and other adjacent territories. Since bugs, and not least, E. prolixus, are frequently used routinely for xenodiagnosis to assist clinical diagnosis of infection in man, and also in experimental or "wild" infections in other vertebrates, evidence for a genetic mechanism in bugs controlling infection rates in groups, and levels of faecal infection in individual bugs could possibly contribute to improved efficiency of xenodiagnosis by selective breeding of highly susceptible bug populations to be used for the purpose. It might also aid interpretation of infections in wild populations of bugs, although many other factors would also be important in nature. Admittedly, infection rates for T. cruzi, whether natural in wild bugs or experimentally derived, as for example in xenodiagnosis, are high for virtually all species adequately documented,

and this is, at the present, a pertinent criticism to the geneticist. Evidence, as cited later, for Mendelian inheritance of resistance in other insects in their vector functions in filariasis or malaria also justified the genetical study being carried out with *H. tritarsus* and *T. tritarsus*.

The results reported in the foregoing pages show that detection and interpretation of the genetic control of bug susceptibility was by no means simple and on the basis of percentage infection rates the data obtained did not support a Mendelian controlling mechanism. On the other hand, analysis of first and second degree statistics derived from quantitative assessment of fecal infections in parental bugs and their progeny, elicited evidence of polygenic control of infection with *T. tritarsus* in the form of response to two-way selection over 3 generations for populations with increased or reduced ability to support infection. In a general way discussing the design, it is worth mentioning briefly the selection procedures adopted and to follow that with discussion of the more important findings of which the conclusions drawn are worthy.

After an infecting blood-meal as fifth instar larvae, *H. tritarsus* adults were given a clean blood-meal and the numbers of trypomastotes excreted estimated by scoring standardly diluted samples of their faeces in terms of the number of trypomastotes per 100 fields. Two parental groups of bugs infected in this way with different strains of *T. tritarsus* (Jeru strain and Strain 7) were scored and selected parents from both groups were mated. The progeny (first generation) were infected, given an infecting feed as fifth instars, and scored when moulted to *T. tritarsus* adults; selected bugs were then again infected, and the process was repeated with the 3 generations of bugs infected with strains of trypomastotes, and again the 3rd generation of bugs infected with *T. tritarsus* Strain 7.

Refractory by Class

Bugs were classed as susceptible to T. cruzi infections if they showed any sign of faecal infection following an infecting blood-meal on mice, or as refractory if no trypanosomes of any form could be detected either in the faeces or in recipient clean mice injected with samples of the faeces. In the parental generation 11.6% of bugs tested were refractory to infection with T. cruzi Peru strain, while 10.4% of bugs infected with Strain 7 were refractory. Alternatively the populations may be considered as having susceptibility rates of 86.4% and 89.6% respectively, which are very close to the figures of 88.4% obtained by Phillips and Bertine (1967) based on the mean of 3 groups of H. triolix, infected with different strains of T. cruzi.

Attempts to increase the proportion of refractory bugs within families by sib-mating refractory or low scoring parents failed in both experimental groups. For example, after the generation of selection the proportion of refractory bugs in F_2 family 18 of the Peru strain group was 18.0%, a figure not significantly different from the proportion of refractory bugs in the parental generation of 11.6% (corrected $\chi^2 = 0.006$, $P = N.S.$) and yet this family had a greater proportion of refractory members than any other F_2 family selected for refractoriness. Similarly, selecting the refractoriness free bugs infected with T. cruzi Strain 7 produced F_2 families with a mean proportion of 18.7% refractory which was not significantly different from the parental generation figure of 10.4% ($\chi^2 = 5.4$, $P = 0.05$). Statistical tests to detect any trend towards an increase in the proportion of refractory bugs as a result of selection were negative whether bugs were infected with Peru strain or Strain 7 of T. cruzi, respectively. Furthermore, F_2 families 11 and 12, which contained the

greatest proportion of refractory bugs, were descended from susceptible parental stock.

Selection for increasing proportions of susceptible bugs infected with Peru strain 1, 1961 gave similar discouraging results, there being no significant effect by the F_2 generation and, moreover, there was no significant difference in proportions of susceptible bugs between the two groups of F_2 families selected for susceptibility and refractoriness. However, families selected for susceptibility to Peru strain 7 over 3 generations did show significant increases in the proportions of susceptible bugs within families. Selected F_2 families had a mean proportion of 37.3% susceptible members which was significantly greater than the 27.6% in the parental generation ($\chi^2 = 4.0$, $P = 0.05-0.01$). A test for linear trends in proportions confirmed that over the 3 generations there was a systematic increase in the proportions of susceptible bugs in these selected families.

The results of these analyses are presented in Table 1. Selection for refractory bugs (as defined in the present work) was clearly unsuccessful. Phillips and Morton (1961, 1962), who have provided the only comparable data increased the proportions of refractory bugs in the F_2 to 41.4 from the parental mean of 24.6 by selecting refractory 1961 families. A survey of the F_2 data of the present experiments shows that high proportions of refractory bugs were obtained in some families of the Strain 7 Infected group; for example, F_2 family 70 had 45.5% refractory bugs (10 out of 22) but subsequent selection and assortative mating from this family resulted in F_3 families 4 and 5 with 5.2% and 2% refractory members respectively. The results obtained from F_2 family 73 of the Strain 7 group are interesting, where selection for refractoriness from this family, which was 5.9% refractory, produced

3 F_2 families F_1 70 and 11 with 5.4%, 41.2% and 46.7% of refractory members respectively. The parents of F_1 family 73 were both susceptible which makes a Mendelian interpretation of the data rather difficult. If susceptibility were controlled by a single Mendelian recessive factor then both the parents of F_1 family 73 would have to be homozygous and could not, therefore, have produced refractory offspring. Alternatively, if refractoriness were the dominant or allele then two susceptible parents of F_1 family 73 must be assumed to have been heterozygous and should have produced a 3:1 ratio of susceptible to refractory progeny. If refractoriness were dominant then F_1 generation parents should then have all been refractory as should the F_2 offspring. This, clearly, was not the case. However, selection for susceptibility did produce significant changes in the proportion of the 2 classes in families infected with Strain 7 trypanosomes which suggests that susceptibility is, at least in part, genetically controlled, but these results give no indication as to the nature of the control. The fact that selection for susceptibility was successful while selection for refractoriness was not, but that a Mendelian explanation for refractoriness were controlled by a recessive allele (a) with a dominant allele (b) for susceptibility, then the failure to select for refractoriness may be explained as a failure to recognize homozygous (aa) bags phenotypically, indeed the test system adopted for the present experiment. This would produce results which do not fit expected Mendelian proportions. Alternatively, it might be that the significant differences in proportions of refractory bags between F_2 families of the Strain 7 group, selected for refractoriness and susceptibility ($\chi^2 = 11.1$, $P < 0.001$ ***) were a type of side effect of changes in the intensity of infection brought about in the 2 groups.

Suppose that the intensity of infection of a bug was reduced to a low level by selection, then the correct phenotypic identification of this bug as refractory, or susceptible, would have been difficult because the method of checking microscopically negative faeces by inoculation into a clean mouse could overlook a very low infection if the immune response of the recipient mouse eliminated the infecting parasites.

In conclusion, a major gene interpretation of low gene frequencies here, that is that susceptibility to L. major infection in M. musculus is controlled by alleles at a single locus, cannot be supported, but neither has such an interpretation been completely excluded by the present work. Further generations of selection and inbreeding, not this being practicable, might have clarified the situation, but this could be doubted as, in a similar selection experiment in M. lauraxii, Macdonald (1962a) produced a susceptible, true-breeding line of the susceptible domestic mouse, in the F_2 generation using inbred strains, and was able to demonstrate that the gene controlling susceptibility to the filicidal agent, Trypanosoma, in the mouse was a dominant recessive factor. Moreover, Macdonald (loc. cit.) was able to increase the proportion of susceptible inbred Form 17 to the parental generation to 85.7% in the F_2 . The present work has failed to demonstrate such dramatic change in the proportions of either susceptible or refractory bugs as a result of selection and inbreeding, but while this does not rule out a major gene control factor, such an hypothesis has not been substantiated.

I turn now to the conclusions drawn by quantitative analysis dealing with levels of infection of trypanosomes in the faecal excrement of individuals M. musculus, domestic, Form 17 of Trypanosoma per individual sample of bug faeces.

Quantitative analysis

The intensity of infection of each infected bug was assessed by counting the number of trypanosomes in 100 fields ($\times 450$) or a 20 μ l sample of faeces of a standard dilution. The resulting scores were transformed to $\log (n + 1)$ to normalize the data for statistical analysis. Changes in family mean and variance following infection and sib-mating were analysed to determine whether intensity of infection was a heritable quantitative character. These analyses were carried out on the scores per 100 fields obtained from the same bugs which were analysed by class as positive or negative. The third analysis was designed to detect differences in intensity of infection between bugs infected with the 2 strains of trypanosome. Although the experiment was not strictly replicated, it is not impossible, if scores of 100000 test the same group of bugs with both strains of trypanosome, the bugs used in the parental generation were randomly chosen from the same colony and may be assumed to be genetically homogeneous. At this parental generation, significant differences were shown in intensity of infection between bugs infected with the 2 different strains; bugs given an infecting feed on T. cruzi Strain 7 had a mean score approximately twice that of bugs fed with the more virulent Peru strain. This result is most surprising since the bugs ingesting the Peru strain (a virulent, lethal strain for mice) were fed on mice with higher levels of parasitaemia than those infected with Strain 7 T. cruzi (a strain of low parasitaemia from which mice could survive). It can be assumed that both groups of bug took approximately the same size blood-meal, therefore Peru strain infected bug must have ingested more trypanosomes. Yet bugs excreted fewer than the Strain 7 group of bugs, an indication that the virulent trypanosomes of the Peru strain, possibly at low numbers, had a

lower potential for producing a correspondingly heavy infection in the vector bug.

There are no comparable quantitative studies of the nature covered in the present work. In the literature, on the intensity of infection of different strains of trypanosome in the bug. True, Nydomen (1965) did report a quantitative difference in the density of trypanosomes in the faeces of triatomines infected with different L. tritici strains; also the effect of trypanosome strain on susceptibility rates (i.e. proportions of infected bugs in a population) of triatomine bugs was studied, and strain differences were recorded (Ovdiuk and Butrim (1967) and Ovdiuk and Simida (1968)). But these studies were not designed to follow the inheritance of quantitative changes in intensity of infection within bugs. Further analyses of F_2 families of the present series again showed highly significant differences in intensity of infection in individual bugs between groups infected with the 2 different strains of trypanosome. Analysis of further generations has not carried out since the effects of selection may have affected variation between strains disproportionately. However, it may be concluded from the results from F_1 families, and from the parental generations that intensity of infection in P. prolixus is affected in part by the genotype of the strain of trypanosome which the bug ingests.

A particularly interesting finding was obtained by analyses of variance of F_1 and F_2 families infected with the 2 strains of trypanosome in that all analyses revealed significant differences in intensity of infection of male and female bugs when tested against the considerable variation between families. Highly significant differences ($P < 0.001$) were also found between sexes in F_2 families of the Birko 1 group. These differences between sexes were constant for the several families of both

lower potential for producing a correspondingly heavy infection in the vector bug.

There are no comparable quantitative studies of the nature covered in the present work, in the literature, on the intensity of infection of different strains of trypanosome in the bug. True, Kyriakou (1968) did report a quantitative difference in the density of trypanosomes in the faeces of triatomines infected with different *T. brucei* strains; also the effect of trypanosome strain on susceptibility rates (i.e., proportions of infected bugs in a population) of triatomine bugs was studied, and striking differences were observed in *Triatoma* and *Recliniterm* (LSD) and *Triatoma* and *Recliniterm* (LSD). But these studies were not designed to follow the inheritance of quantitative changes in intensity of infection within *Triatoma*. Further analysis of F_2 families of the present series again showed highly significant differences in intensity of infection in individual bugs between groups infected with the 2 different strains of trypanosome. Analysis of previous generations must not be ruled out since the effects of selection may have affected variation between strains disproportionately. However, it may be concluded from the results from F_2 families, and from the parental generations that intensity of infection in *M. proligus* is affected in part by the genotype of the strain of trypanosome which the bug ingests.

A particularly interesting finding was obtained by analyses of variance of F_2 and F_3 families infected with the 2 strains of trypanosome in that all analyses revealed significant differences in intensity of infection of male and female bugs when tested against the considerable variation between families. Highly significant differences ($P < 0.001$ ***) were also found between sexes in F_2 families of the Strain 7 group. These differences between sexes were constant for the several families of each

group tested since no significant sex X family interactions were detected in the analyses. It was thought with justifiable reference to these sex differences, that they may have been attributable to differences in the volume of blood taken at an infecting feed at their fifth instar blood-meal or to differences between sexes in rate of excretion when tested as adults. Further experiments showed that there were no significant differences in blood-meal size between male and female fifth instar *K. proxima* larvae; differences in numbers of trypanosomes ingested could not therefore, have produced the sex differences. Also, by weighing faeces collected within 1 hour of feeding (see Table 1), it was found (see Table 2) that adult females took longer to defecate, excreted a greater mass of faeces and excreted a greater proportion of their blood-meal within 1 hour of feeding. A reasonable expectation would be that these findings, that female excreta would have been greater than male excreta but the results showed the female to be the group, whose excreta being significantly greater than female excreta throughout the selection experiment. The conclusion follows that intensity of selection for individual bugs is influenced by the sex of the bug ingesting trypanosomes regardless of the strain of *K. proxima*. Thus, sex is a genetically determined character in bugs, these results provide good evidence for the involvement of bug genotype in determining the numbers of trypanosomes which develop in a bug's gut after an infecting feed.

The analysis of F_2 families infected with strain 7 showed a further interesting result in which there was a significant sex X selection line interaction when tested against the error term. Differences between sexes were greater in families selected for high score than in those selected for low score. This result may have been produced by sex-linkage of genes controlling susceptibility or, alternatively, may

have been the result of a failure in scaling by which the log transformation did not fully remove the positive skewness evident in the frequency distribution of the raw data. The precise nature of the genetic control of sex-differences in intensity of infection could not be determined from the present work. It may be that the control of susceptibility is sex-linked but the significant sex-differences may have been produced by sex-linked genes. Williams and Barton (1967) found no significant difference in infection rate between sexes of R. prolixus experimentally infected from mice and Ansidei et al (1973) failed to detect a difference in susceptibility rate between sexes of T. trypomastix. However, neither of these studies was based on precise quantitative estimates of the intensity of infection, and are not strictly comparable with the present work.

A constant feature of the analyses was the significant differences between families. These between-family differences were detected at the analysis of trypanosome strain differences, indicating that family differences were to be found in both groups of bugs infected with different strains. In the context of sex differences, no between-family item was found to be highly significant in all the analyses of variance of bug scores for the F_1 and F_2 generations and for both strains of trypanosome used. Furthermore, tests for sex \times family interaction indicate parent specific, individual family differences were independent of the significant sex differences. These significant between family differences are of great genetic importance for they signify resemblance between related individual bugs for a quantitative character (intensity of infection) since significant differences between several families must reflect the similarity of individuals in any one family. The resemblance between phenotypes in

one of the basic characteristics displayed by metric characters (Falconer, 1967) and these differences in variance are a strong indication that intensity of infection in K. bollux is polygenically controlled.

The analysis of F_2 females infected with T. viris Strain 7 (Table 3.47) showed only borderline significant differences between families ($P = 0.05-0.10$) and was in contrast to the analysis of male F_2 scores of the same families which showed highly significant differences between families (Table 3.46; $P = 0.001$). This difference may be explained by the fact that the item tested in Table 3.47 was variation between female families within selection lines and it is to be expected that after selection and breeding by F_2 , variation between families within a selection line would be reduced. This result may be taken as an indication of the effectiveness of the selection procedure for females which may have been reaching the limits of their selection potential, at which point it would be expected that family means of a particular selection line would be in close agreement.

The degree of resemblance between relatives is a property of a polygenic character which can be quantified by experimental means or by analysing the results of a selection programme. The degree of resemblance between relatives provides an estimate of the amount of additive variance, and it is the proportionate amount of additive variance which determines the breeding value or heritability of a polygenic character. The heritability has great predictive value to breeders and is a most important property of a metric character, the greater the proportion of the heritable variation which is additive and stable, the greater will be the effectiveness of selection. The heritability of intensity of infection was estimated, in the

present work, from the path of response to the selection differential (S) from a two-way selection programme applied to B. prolixus infected with T. cruzi strain 7. Two lines were derived from the parental population of bugs selected for high and low levels of intensity of infection over 3 generations. In the F_2 generation, highly significant differences were found between bugs of the 2 selection lines of both sexes ($P < 0.001$ ***). Furthermore, when the F_2 family with the lowest mean score from the high line was compared with the family with the highest mean score from the low line, differences between these 2 families were highly significant ($P < 0.001$ ***) when compared with the variation between high scoring families. Thus, selection from the parental generation of randomly selected bugs produced in the F_2 two discrete populations with significantly different trends, as a legitimate study of heredity is hereditary. The realized heritability for the character was determined from the regression of response on selection differential for both selection lines. The regression lines for both lines were highly significant ($P < 0.001$ ***) so that the slopes of the lines provided sound estimates of the heritability of this character. The response to selection measured from the divergence of 2 reciprocal lines, taken as twice that of the non-operating, produced an estimate of 5.0 for the heritability of intensified infection in B. prolixus under the experimental conditions of this work.

Reverse selection from F_2 families infected with T. cruzi strain 7 did not, however, produce F_2 families with mean scores significantly different from the families selected throughout the high scores. This result is, perhaps, not surprising after only one generation of reverse selection. However, when the regression of response to selection differential for the reverse selected high line was carried out, the result

was a negative slope ($b = 0.1181$) and the regression was significant, indicating that there was no longer a linear upward trend in mean score after reverse selection. Taking these two findings together, it may be concluded that reverse selection had a small effect on family scores but this effect was not large enough to produce significant differences when compared with the variation within families.

When the regressions of response on selection differential for high and low lines were further considered it was clear that the response to selection pressure was not equal in both directions (Fig. 4b). This asymmetry of response is a common feature of two-way selection programmes with animals and has been attributed (Falconer, 1960) to two main causes: inbreeding depression and directional dominance. It is known that inbreeding can lead to a reduction in the mean phenotypic value of a quantitative character which is connected with fitness - such as reproductive capacity (Falconer, 1961). Correlated responses with low heritability are often related to fitness; for example, the amount of white spotting in Friesian cattle has a very high heritability of .75 (Briquet and Lush, 1947) while conception rate (which is fitness related) in the same cattle has a very low heritability of only .16 (Wentworth, 1967). The low heritability found in the present work for intensity of infection (.5%) suggests that this character may be related to fitness and therefore subject to inbreeding depression. Populations which show inbreeding depression will respond more rapidly to downward selection than to upward selection and, since the asymmetry of response demonstrated for *M. paratuberculosis* showed downward selection to be twice as effective as upward selection, this may further indicate that intensity of infection is related to fitness. Alternatively, it could be argued that the genes controlling intensity of infection are merely linked to

those of another character more directly concerned with fitness or, it may be, that in selecting for trypanosome density in bug hosts an unconscious selection of the fittest individuals was taking place. It is important to consider, therefore, what direct effect density of a trypanosome infection in the gut of a bug could have on the fitness of a bug. Hare (1971) has suggested that mammalian trypanosomes have evolved from the monogenic trypanosomatid parasitic flagellates of two-blood-sucking insects, and that bugs infected with *blutauschillia* acquired the blood-sucking habit, demonstrated at low rates of the bite and the trypanosomes of their gut entered the mammal where eventually they established themselves as *hematophagous*, the bugs being reduced to intermediate host status. Whether this hypothesis is true or not, it is known that the relationship between *T. evansi* and the triatomines is not one of *obligate* parasitism and, if the trypanosomes were once solely endoparasites of bugs, it is to be expected that bugs would have evolved genotypes specifically adapted to cope with the *parasitoid* relationship, and they will surely be related to the fitness of the bug.

Falconer (1967) also states that if the genes that increase a metric character are dominant over their alleles responsible for a reduction in its value, then *inbreeding* will result in a reduction in the population mean for the character, i.e. a change in the direction of the more *recessive* alleles. Thus, it would be expected that a character which showed inbreeding depression would respond more rapidly to *downward* selection, in which case the results of the present work suggest intensity of infection may be related to fitness or *biological* and also that the genes for increasing susceptibility could be dominant to those for *infectiousness* and decreasing intensity of infection.

Knowledge of genetic mechanisms controlling the susceptibility of insect vectors to infection is limited to mosquitoes and their malarial and filarial parasites. Briefly, Huff (1929) was the first to suggest that susceptibility of a vector was genetically controlled from his study of susceptibility of Culex pipiens to Plasmodium latrans. Further experiments by others were subsequently done with various mosquito vector-malaria associations but the mode of inheritance remained obscure until recently when Kilama and Craig (1969) demonstrated that susceptibility of Anopheles gambiae to infection with Plasmodium falciparum was controlled by a pair of autosomal recessive genes.

The genetic control of susceptibility of the mosquito Anopheles gambiae to infection with the filarial nematode Brugia malayi was first investigated by Rouboud et al. (1936) who found variation in the susceptibility rate of strains of mosquito from different geographic regions. Rouboud (1936) suggested that susceptibility of the mosquito Anopheles gambiae to filarial infection with Brugia malayi was controlled by a sex-linked recessive gene. Recently Jelleke (1973) and McNamee et al. (1974) extended these studies, showing that the susceptibility of An. gambiae to another filarial worm, Wuchereria bancrofti, is also controlled by a single sex-linked recessive gene.

It is notable that this work on susceptibility of mosquitoes to parasites which they transmit has so far revealed only major gene control mechanisms. There have been attempts to quantify mosquito susceptibility in terms of infection levels of individual insects but these have not produced clear-cut results. For example, Brown (1964) was unable to produce parent-offspring regressions for many years. However, it is interesting that selection experiments with mosquito filarial systems have often failed to produce completely refractory or completely susceptible

stocks of mosquitoes even after many generations of inbreeding. For example, Macdonald (1962a) selected a strain of An. stephensi susceptible to infection with Pr. m. m. m. for 15 generations at which point the susceptibility rate was still only 84.8%. Macdonald et al. (1974) attributed the variation in susceptibility rate of selected populations of An. stephensi infected with Pr. m. m. m. to the fact that the male genotype cannot be determined. It may be that intensity of infection in mosquitoes is controlled, in part, polygenically and this could influence the interpretation of data concerned, at present, only with dividing populations into - classes, refractory and susceptible.

The present work has demonstrated that intensity of infection with Pr. m. m. m. in An. stephensi is a quantitative character with low heritability and that this character is sex-linked, or sex-limited. It is further suggested that genes for increasing intensity of infection are dominant to those for decreasing intensity. This experiment has failed to demonstrate that susceptibility to infection with Pr. m. m. m. in An. stephensi is controlled by major genes, but further generations of selection would be necessary to clarify this point. Mather and Jinks (1971), in their work on biometrical genetics, suggest that with the possible exception of antigenic traits, all the characters of an organism are subject to both continuous and discontinuous variation as a result of minor gene and polygenic differences. It is, therefore, unwise to rule out the possibility that a major gene mechanism may be involved in the susceptibility of An. stephensi to Pr. m. m. m. infection, as well as the environmental component control of intensity of infection. Finally, the question of xenodiagnosis must be mentioned. This experiment has shown that it is possible to increase the intensity of infection in An. stephensi by selection. For the purposes of xenodiagnosis

it would be desirable to breed and maintain in standard laboratory culture, bugs which, on feeding on patients with very low levels of infection, would be so highly susceptible to multiplication of the few trypanosomes ingested that they would be detected readily within a month or so in the bug faeces. Since it has been shown that the heritability of this character is low, the best method of producing such a population of bugs would be by using family rather than individual selection (Falconer, 1967). The present work indicates that, having established an inbred line of bugs with high levels of intensity of infection, only male bugs should be used for xenodiagnosis (usually, late pre-adult larvae are used, but it is a mistake to use fifth instar bugs - usually, 1000-1500 are used from one patient). The prospects are that, with such a line, the sensitivity of this diagnostic test would be considerably improved, and with the added benefit of using fewer bugs per patient, by no means a negligible matter from the point of view of the psychological effect some patients feel with this technique, and the inevitable problem of maintaining adequate supplies of bugs.

P A R T F O U R

THE INHERITANCE OF RADIATION INDUCED SEMI-STERILITY

IN NECROMUS PROLEPS

STERILIZATION

In recent years several genetic methods have been proposed for the control of insect pests, both of agricultural and public health importance. The subject was recently reviewed by Knippling (1977). Because of its demonstrated efficacy, the sterile male technique has become the most widely known and studied of these genetic control methods. The sterile male technique, first advocated as a practical reality by Knippling (1955), involves sterilizing males by irradiation or chemosterilization and releasing them, usually in vast numbers, into the wild population to mate with and inseminate normal wild females with resultant sterility of those eggs. It was found that low doses of mutagen required to induce the essential chromosomal damage in the sperm of diploid male insects was of minimal importance, the aim being to give a dose which would not affect male competitiveness yet would ensure that transmitted sperm would transmit recessive mutations which at fertilization would result in offspring lacking functional chromosomes. First working Miller (1947), followed by a series of complex experiments with the parthenogenetic and *phlebotomus papatasi* to result from chromosomal breakages rather than from point mutations (Smith and von Borstel, 1972). Knippling's mathematical model of the population dynamics of sterile male release (Knippling, 1961) was put on trial in the field and the first successful results were achieved in Curacao when the screwworm fly *Cooperia hominivorax* was eradicated from the island by releasing sterilized males (Knippling et al., 1961). Following this success, the sterile male technique was extended to the mainland of the United States and the screwworm fly was eliminated from the state of Texas (Knippling, 1961). From after this success, the method was extended to the States bordering on Mexico,

where a barrier was set up along the border. This biological barrier has involved releasing one thousand million sterile flies per year along the Mexico-Texas border (Bushland, 1971). Recently this control programme has apparently suffered a setback, a recrudescence of large numbers of "strikes" by screw-worms, particularly in Southern U.S. territory contiguous with Mexico being reported in *Nature* (1973), and attributed to the reduced fitness of the factory-bred flies in competition with the wild-type. However, following the initial success of the screwworm fly control programme, the sterile male technique has been tried with several other insect pests, especially those of economic value. One problem for several populations to eradicate since re-invasion by neighbouring populations is precluded; for example, the onion fly *Delia floralis* has been eliminated from the island of Rota in the Pacific Ocean by sterile male release (Steiner et al., 1965).

The success of sterile male control programmes stimulated interest in alternative forms of genetic manipulation and, in particular, in the release of partially sterile males or fully fertile males carrying inherited characters which would affect not only the first but also subsequent generations of the species. Several authors have been proposed for the production of insects with inherited sterility factors including chromosome translocations, recessive lethal mutations, chromosomal mutations, and meiotic drive which would produce sex ratio distortions. The idea of using translocations to control insect populations was proposed initially by Bernabrovsky (194) and later by Curtis (1968). A chromosomal translocation arises when two non-homologous chromosomes in the same cell are broken and the fragments are interchanged. ~~The process of interchange production is~~ The process of interchange production is illustrated in Fig. 4.1 which shows that when meiosis occurs in the heterozygote formed by the union of an egg with a sperm carrying a translocation, a cross configuration must be assumed at

reduplication in order that the chromosomes may pair. If the cross is formed in each of the arms of such a cross, the bivalents form a ring at first metaphase (Fig. 4.2a) and if in the one in which this metaphase ring disjoins (Fig. 4.2b) which produces sterile gametes are produced. Balanced gametes only result from the so-called "alternate" disjunction when alternate centromeres of the metaphase ring go to the same pole at anaphase which involves a twisting of the ring multiple. If adjacent centromeres proceed to the same pole then the resultant gametes will be genetically imbalanced because of duplication and deficiencies of chromosomes (Fig. 4.2b). Of the balanced gametes produced by alternate orientation, half are wild-type and half carry the translocation and can pass it on to the next generation in which a heterozygote would again produce balanced and sterile gametes. Clearly, the mode of orientation of the ring multiple is critical in determining the fertility of translocation heterozygotes. Although 6 possible genetic types of gamete may be produced by a ring multiple (Fig. 4.3) they are not in practice produced with equal frequency and it has been shown that a single translocation will usually result in producing a heterozygote with 50% fertility i.e. producing a 1:1 ratio of balanced to unbalanced gametes (Curtis, 1968). This apparent anomaly is a result of the non-random orientation of ring multiples; John and Lewis (1965) state that the expectation is 1:1 alternate to adjacent orientation in organisms with a single centromere if random behaviour of the two co-orientated adjacent centromeres is assumed, hence the 50% sterility of the translocation heterozygote.

Serebrovsky (1940) and Curtis (1968) further proposed that if translocation heterozygotes were inbred, insects homozygous for the translocation could be produced which on mating with the wild-type would again yield semi-sterile offspring; they further suggested that this

translocation homozygote would be fully fertile since it would be genetically balanced. This hypothesis has been substantiated for some insects; for example, Laven et al. (1971) produced fully fertile Drosophila homozygotes for an autosomal translocation. Unfortunately, translocation homozygotes may suffer from reduced fitness in competition with the wild-type insect population (Curtis et al., 1977). Bearing this loss of fitness in mind, and also considering the very low yield of viable translocation homozygotes which is normally achieved by irradiation treatment of normal stock insects, McDonald and Kai (1971) suggested that translocation heterozygotes be released in control programs. This method could also be used for species which cannot be mass produced, when wild caught insects could be partially sterilized thereby introducing translocations into the population (Laven et al., 1971). It has also been suggested (Curtis and Robinson, 1971; Whittam, 1971) that multiple translocations would be more desirable for pest control programs than single interchanges as they would yield a higher level of sterility. A disadvantage of controlling insect populations by releasing semi-sterile males could be that, instead of eradicating a pest, the population would be simply replaced by the translocation-carrying population. This could, however, prove advantageous if the translocation stock was also carrying desirable genetic linkages. Curtis (1968), for example, has suggested that in this way a wild-type population could be replaced by a translocation stock carrying genes for susceptibility to insecticides or refractoriness to parasites.

Biokinetic types. Schrader (1936) first noted that different forms of centromere were to be found in chromosomes of different species; firstly, a sharply localized type and secondly, a diffuse type. The diffuse type of centromere was later called holokinetic (Huxley, 1953).

holokinetetic (or holocentric) chromosomes show no primary constriction and the spindle fibers appear to be attached along the entire length of the chromosomes. Organisms with holokinetetic chromosomes are sporadically distributed throughout the animal and plant kingdoms. In the animal kingdom, holokinetetic species are limited to the arthropods and are most common amongst the insects (John, *et al.*, 1965), and holokinetetic chromosomes are known to be typical of the hemiptera (Hugues-Schrader and Schrader, 1961). It is important to appreciate that a diffuse centromere imparts certain characteristics to a chromosome. For example, at mitotic anaphase holokinetetic chromosomes proceed to the poles with their long axes perpendicular to the long axis of the spindle (Schrader, 1953). It was on these behavioral characteristics that holokinetetic chromosomes were recognized. In many organisms and, subsequently, experimental proof of their structure was obtained. Hugues-Schrader and Riss (1941), were able to show that X-ray induced fragments of cockle chromosomes, behaved normally at mitosis, each fragment, regardless of its size, dividing and passing to the poles at anaphase through many cell divisions. The peculiarities of the holokinetetic system later proved to be useful in insect control programmes involving partially sterile males. It was observed by Provencio (1962) that male coding moths (*Lappogryllus pennsylvanicus*) given a sub-sterilizing dose of radiation and then mated to normal females resulted in reduced numbers of F_1 progeny, the majority of which were males with very low fertility. This delayed effect of irradiation in lepidoptera was confirmed by other workers including North and Holt (1966a) for cabbage looper (*Trichoplusia ni*) and Walker and Quintana (1966) for the sugarcane borer (*Plutania maculipennis*), the level of sterility being greater for both species in F_1 progeny than in the

treated male parents. The use of sub-sterilizing doses of radiation in these experiments had not been accidental since it was already known that the Lepidoptera were highly resistant to induced reduction in fertility by irradiation; for example, 30-40 K rad had been required to completely sterilize codling moths in an experiment carried out by Proverbs and Newton (1963). Such high dosage produced considerable damage and rendered the males less competitive so that male cabbage loopers given high doses of gamma-rays failed to transfer any sperm when mating with normal females (North and Holt, 1968b). The explanation for this high degree of radio-resistance encountered in lepidoptera, and for the phenomenon of delayed sterility of males given sub-sterilizing doses was provided by Bauer (1967) working with *Pieris brassicae*. He compared the observed rate of viable translocations in this butterfly, after irradiation, with the expected rate for a kinetochore region and, from this, proposed that, in a holokinetochore region, probably due to a single translocation, could not occur at all or only exceptionally for the following reason: Following a break in two non-homologous chromosomes in a monokinetoc animal there are two possible results; firstly, the proximal broken ends can unite to produce a dicentric chromosome while the distal ends join to form a fragment or, secondly, a symmetrical reciprocal translocation can be produced as shown in Fig. 4.1. Normally, dicentric chromosome production will lead to genetic duplication-deficiencies due to bridge formation at anaphase and to the loss of the acentric fragments, but Bauer (loc. cit.) proposed that in holokinetoc species dicentric chromosomes could not be formed. Indeed, his experimental results confirmed this hypothesis for he obtained a much greater proportion of viable translocations than would be expected with a monokinetoc organism. In fact, Bauer obtained 2%

zygote lethality in this experiment which was much higher than he expected from his hypothesis and he attributed these deaths to gene mutation or extra-nuclear radiation effects, maintaining that they could not have been caused by small chromosomal fragments.

North and Molt (1968b) confirmed Bauer's hypothesis when they found, by microscopical examination of large numbers of translocations in the spermatocytes of F_1 male cabbage loopers whose male parents had been given sub-sterilizing doses of radiation, some translocations, which they suggested, were the result of 'dicentric' chromosomes behaving as reciprocal translocations in this holokinetic species. North and Molt (loc. cit.) provided further cytogenetic explanation for the sterility of F_1 males being greater than that of their irradiated parents, by stating that reciprocal translocations could result in the formation of quadrivalent-chromosome complexes, depending on the mode of orientation of the multiples at metaphase, so that when F_1 spores were used in fertilisation they could be lethal to the embryo, the degree of F_1 sterility depending on the number of interchanges induced by the radiation.

In addition to the production of reciprocal translocations, diffuse centromeres also ensure that chromosomal fragments which would normally be lost in monocentric organisms, are mitotically stable in holokinetic species. LeThomas and Desregulier (1962) working with the holokinetic milkweed bug, Oncopeltus fasciatus, showed that X-ray induced fragments were not only mitotically stable but were also meiotically stable and could be transmitted through 3 generations of outcrosses to normal females. Furthermore, LeThomas et al. (1961) demonstrated that a single chromosomal fragment could lower the fertility of O. fasciatus appreciably, the degree of sterility depending on the size of the

fragment and its pairing behaviour so that 3 fragments produced almost total sterility while random segregation of a single fragment produced 50% sterility. Tomaz-Nunes's (1967) hypothesis that chromosomal fragments were not responsible for embryonic deaths was shown to be incorrect. Kripling (1970) devised mathematical models to estimate the effect of the release of partially sterile Lepidoptera on wild populations and concluded that in mass releases, such males would be more effective in suppressing pest populations than completely sterile males. It is clear from all this work with Lepidoptera that centromere structure can play an important part in determining the radiosensitivity of an insect and in deciding the correct approach to controlling a wild population.

The control of Triatomines (Hemiptera, Reduviidae) is of great interest because of their importance as vectors of 'kissing' disease and some work has been carried out to investigate ways of sterilizing these pests. Tomaz-Nunes et al. (1962, 1964) and Baldwin and Chant (1963) have studied the effect of irradiation on Triatoma prolixum and have produced dose-response curves with respect to fertility. Tomaz-Nunes et al. (1964) using γ -irradiation and Baldwin and Chant (1963) using X-rays found that 20 K rad were required to sterilize completely adult T. prolixum, this dose being comparable to that required to sterilize Lepidoptera (North and Molt, 1968b). Tomaz-Nunes et al. (1964) also studied the effect of irradiation on longevity and competitiveness and found that at 5 K rad irradiated males were as competitive as normal males and produced viable sperm, while their longevity was not reduced significantly. Baldwin and Chant (1971) showed that by irradiating T. prolixum in an atmosphere of nitrogen the competitiveness of males given sterilizing doses of X-rays was improved over controls irradiated in air. Sterility studies have also been carried out with triatomine

bugs using alternative methods to irradiation. Chou, et al. (1972) investigated the effect of the chemosterilant metopa on M. prolixus. Trietominae have, however, not received the attention that other tsetse vectors of disease have, in this study, except, particularly in terms of the genetic and cytogenetic mechanisms involved in irradiative sterilization.

Since the Trietominae are Hemiptera, they are of special interest cytogenetically because, as with other reduviid bugs, chromosomes have been assumed to be holokinetic (Ooshima, 1963). With regard to chromosome structure, that of T. tritoma seemed to be atypical because it had a diffuse centromere but found that the kinetochore was apparently absent from achromatic chromosomes. George and Hilde (1971) in an ultrastructural study of Protoparva fuscipes, also showed that kinetochore plates were present along most of the length of mitotic chromosomes while, in the achromatic chromosomes, they could find no kinetochore plates but, found spindle microtubules inserted along the length of the chromosomes which were thought to be responsible for chromosomal movements during meiosis. They suggest that in holokinetic species, suppression of kinetochore activity is essential during meiosis to facilitate chiasma terminalization.

The present study is concerned with the inheritance of radiation-induced partial sterility in M. prolixus. Since oligospermatogenic studies (Buck, 1967) have indicated that M. prolixus has irradiation hypersensitivity, it could be that induced partial sterility in males would be preferable to completely sterile males as a genetic mechanism for control by sterile male release, as suggested by Gomez-Ibanez et al. (1964) and Baldwin and Chant (1970), especially as these workers also showed that small, sterilizing doses of radiation seriously impairs the sexual fitness of M. prolixus males.

MATERIALS AND METHODS

Materials

M. profligator randomly selected from the colony kept at L.S.H. and T.M. were used for this experiment. Details of the history of this colony have already been given.

Methods

All bugs in this experiment were contained in 2" x 1½" glass flat-bottomed tubes covered with fine mesh nylon gauze held in place by adhesive tape. The tubes were kept in incubators maintained at a constant temperature of 27°C and approximately 70% R.H. throughout on top-eared rabbits.

The radiation source was a ⁶⁰Co gamma emitter which gave a dose rate of 2.5 K rad per minute, the facilities being kindly provided by the Department of Physics, Middlesex Hospital, London.

To determine the radiation dose required to produce semi-sterile males, a dose-response curve for fertility was constructed from a preliminary experiment by irradiating adult males, 4 days after molting to adults, in groups of 8 with doses ranging from 0 to 20 K rad in 1 K rad stages. Each male was then, independently paired immediately with a normal female of the same age and seven days later the paired bugs were fed for the first time. They were subsequently fed again every 14 day intervals. The eggs laid after each of the 4 blood-meals were collected and scored for hatchability. Simultaneously, an unexposed control group of bugs were similarly paired and reared and their fertility measured by scoring their eggs in bunches. Both males and females of these series had been fed on rabbit as fifth instars on the same day and had been sexed and separated at this non-mating larval stage so that mating was precisely timed to begin when the paired sexes were set up later in the adult stage, as above.

Since the protocol for the proposed experiment demanded that males be killed for cytogenetic examination, a control experiment was first run to determine the effect on female fertility of removing the male at different time intervals after mating. For this purpose, 2 groups of bugs, each consisting of 7 pairs of males and females, were fed 3 days after moulting to adults and then the pairs were at 14 day intervals, the males being removed from the first group 7 days after the first blood-meal and from the second group 7 days after the second blood-meal. The fertility of the 2 groups was then compared for the 3 blood-meals.

For the main experiment in which the effect of parental irradiation on subsequent reproduction was investigated, 2nd and 3rd instar nymphs were fed on the same day in fifth larval instar, reared then and isolated one per tube. Four days after moulting to adults, 4 males were irradiated, plus 4 together in a separate plastic container, with 6 K rad γ -rays. The males were then immediately paired with the normal females which had been irradiated at the fifth instar, and which were of the same age and nutritional state. These 8 pairs were fed 5 days post-irradiation and subsequently given 3 further meals at 14 day intervals. Mated 1st generation female specimens were fed 4 separate batches, each being provided in a separate and separate and triplicate, the numbers of eggs produced being dependent on the time of the blood-meal (Buxton, 1930; Island et al., 1964). The eggs produced from each of the 4 blood-meals given to the females in this experiment were collected and scored for hatchability. The 1st larvae hatching from these 4 batches of eggs were labelled and kept as a separate batch throughout the experiment and, on reaching the fifth instar, they were sexed and placed individually in separate labelled tubes. On moulting to adults, one normal male, plus or minus, of the same age and placed

In the tube with each P_1 adult according to sex. Eighty-six of these backcross type matings were made, 51 of which were of the form $P_1 \sigma \times$ normal \varnothing and 35 were $P_1 \varnothing \times$ normal σ . Each of these pairs was given 3 blood-meals at 14 day intervals and the eggs laid were collected and scored for hatchability. Seven days after the first blood-meal all P_1 males were removed from the tubes, their testes were dissected out in saline and stored in 3:1 absolute alcohol:acetic acid fixative at 4°C .

For the next generation of matings, parents were chosen only from those $P_1 \times$ normal matings which were of low fertility; the isolated progeny, as well as their parents, were sexed, sexed and isolated up fifth instar to avoid unwanted matings. From 16 such families were chosen and bugs from these families were reared to adults and mated with normal males of approximately the same age.

Twenty-eight of these $P_1 \times$ normal matings were made from among the 16 selected P_1 families. P_1 is used in this experiment merely to denote bugs reared from $P_1 \times$ normal outcrosses and does not imply genetic heterozygosity, 15 of which were of the form $P_1 \sigma \times$ normal \varnothing and 2 of which were $P_1 \varnothing \times$ normal σ . These 28 pairs were each given 3 blood-meals at 14 day intervals and, again, eggs produced were scored for hatchability. Seven days after the fifth blood-meal males were removed, their testes dissected out and stored in fixative. Although only 16 P_1 families were chosen for further breeding, all viable P_1 bugs from all the $P_1 \times$ normal crosses were added to the next stage and although not mated, testes were dissected from the males for cytogenetic examination. The progeny of these P_1 bugs descended from the first egg batch laid by the parental generation were reared to adult, testes dissected from the males and stored in fixative, as

lesions. An assessment was made of the fertility of these ^{60}Co preparations.

Lacto-acetic-orcein squash preparations were made from the fixed testes and a postoperative assessment made of any cytological abnormalities. These cytological examinations were made prior to the analysis of fertility data for each generation and were therefore unbiased by fertility estimates.

RESULTS

Fertility studies

The fertility data for males given increasing doses of X-rays are shown in Table 4.1 together with the control fertility calculated from 10 untreated pairings. All fertility results in this experiment have been calculated as corrected fertility for sterility, this being computed using Abbott's correction (1941), 1961 such that:

$$\text{Corrected fertility} = \frac{\text{Experimental fertility}}{\text{Control fertility}} \times 100$$

A dose-response curve was constructed from the data given in Table 4.1 and this curve is shown in Fig. 4.1 in which, for comparative purposes, dose-response curves constructed from the data of Gammeter et al. (1964) and Baldwin and Shaver (1963) have also been drawn. It is noticeable that the curve constructed from the data of Gammeter et al. (1964) has a very similar slope to that produced in the present work (labelled I.M.), while the curve constructed from Baldwin and Shaver's (1963) data has a slightly different form, although the same species was used in all experiments. These differences may be due to the use of different radiation sources with different dosimetry. Gammeter et al. (1964) used Co^{60} at a dose-rate of 0.075 rad per minute. Baldwin and Shaver (1963) used X-rays at a dose-rate of 1 k rad per

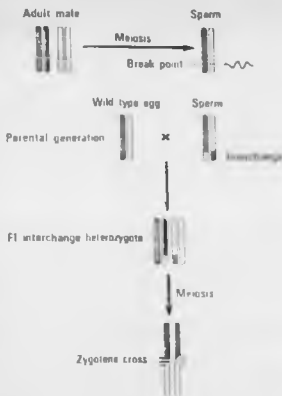


Figure 1. Formation of zygote cross in holokinetetic insects.

RING OF 4 ASSOCIATION

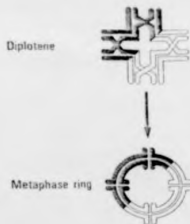


Figure 4.2 (a).

Formation of ring of 4 association in Interchange heterozygotes.

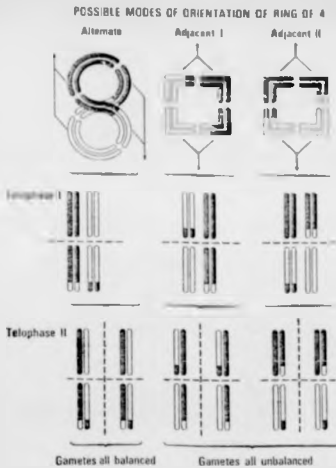


Figure 4.7 (b).

Possible modes of orientation of a ring of 4 association.

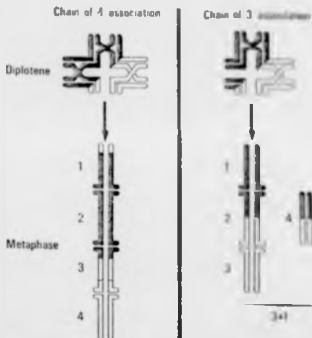


Figure 2-3-1. Formation of chain associations by intercalary heterozygotes.

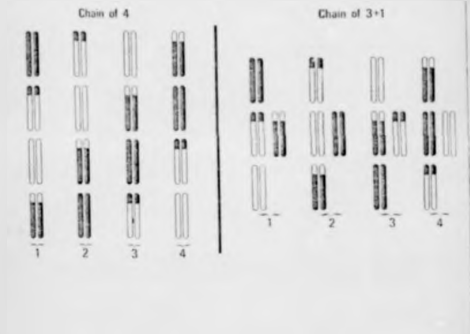


Figure 5.3.10. Possible arrangements of four bivalents, dependent on positions of chiasmata.

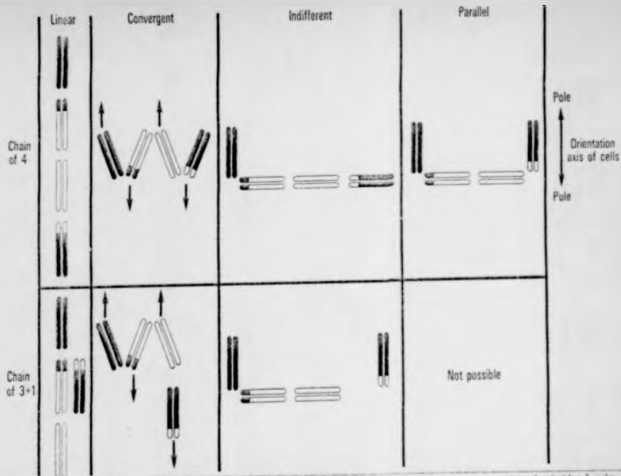


Figure 4.4. Orientation of chain multiples, considering a single association. Only convergent orientation leads to the production of balanced gametes.

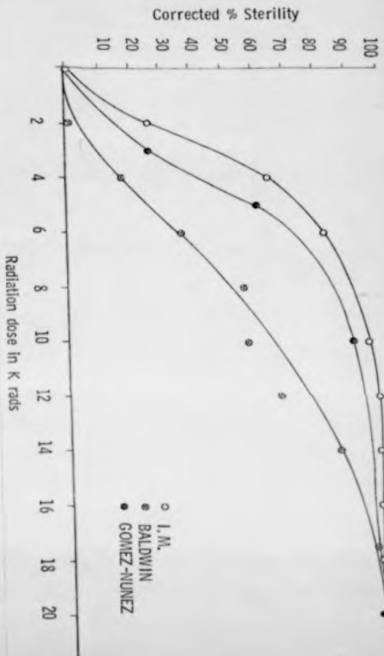


Figure 4.5. Comparison of dose-response curves for *Es. prolinus* compiled from the data of Gomez-Nunez et al. (1964), Baldwin and Shaw (1963) and from the present work (Table 1).

minute. The present work used 10 days at a dose-rate of 2.5 K rad per minute. These qualitative and quantitative differences in radiation applied, probably account for the much smaller dosage required to produce complete sterility in the present experiment (12 K rad) than that needed by Balaban and Gomer (1964) to produce complete sterility (20 K rad).

From this dose-response data, it was decided that 2 K rad should be used as the experimental dosage since this dose produced a high level of sterility (83.16%) and, also, Gomer-Hume et al. (1964) had shown that such a dose did not appreciably affect the mating vigour of male B. pallens. At this point, the cytogenetic effects of such a dose were unknown so that this dosage, per se, was, with particular respect to chromosomal damage, essentially arbitrary.

The preliminary check on time effects on ejaculation and insemination efficiency of normal paired bugs was made as follows: Removing normal males from normal females at 7 days after the first meal (Group 1) or, leaving pairs together for a further 14 days and removing them 7 days after their second blood-meal (Group 2). Using 2 lots of 2 pairs of B. pallens reared on blighty whits. The results of this experiment are shown in Table 4.2. A χ^2 test for proportions, applied to the 2 sets of results, showed that there was no significant difference in fertility between the 2 groups ($\chi^2 = 1.24$, $P = N.S.$), so that removal of the male as early as 7 days after pairing clearly had no effect on hatchability of eggs laid after subsequent blood-meals by the female over a period of 35 days. The data from these two groups were pooled to give a figure of 91.88% for the fertility of 14 pairs of bugs and this was used as the control value for Abbott corrections for subsequent experiments which involved the removal of males after mating.

Table 1.1. Dose-response fertility data for H. prolixa males

irradiated with gamma rays and mated to normal females.

Data pooled for eggs laid after each of three blood-meals.

Dose in K rad.	Number of pairs	Number of egg- laid	Number of eggs hatched	Corrected % sterility
0	4	198	198	0.00
2	4	182	130	26.56
4	4	81	56	50.00
6	4	146	23	83.16
8	4	113	0	100.00
10	4	247	7	96.97
12	4	166	0	100.00
14	4	62	0	100.00
16	4	102	0	100.00
18	4	109	0	100.00
20	4	40	0	100.00

Table 4.2. The effect on fertility of semi-long term *Py. pallidus* from females at different times after mating. The numbers on eggs laid are the totals for 3 blood-meals given at 14 day intervals.

Group	No. pairs	No. eggs laid	No. eggs hatched	% fertility	X ²	P
1	5	507	460	90.73		
2	4	516	324	62.81	14.4	0.001
Total	14	1023	784	76.63		

Group 1 - males removed 7 days after first blood-meal (8 7 days paired)

Group 2 - males removed 7 days after second blood-meal (1 14 days paired)

Having decided on a dosage of 6 K rad, the parental generation of 40 pairs *P. pallidus* were irradiated and each paired with a control female. The hatchability of the eggs laid by these 40 females after 4 blood-meals are shown in Table 4.3, showing that with a dose of 6 K rad the mean fertility of the 20 surviving pairs was 23.0% (all 20 pairs mated and 1 female bug died during the course of this experiment and the data from these matings are not included in the calculations of means). Two pairs had zero fertility but it is important to note that 28 of the 30 surviving pairs were semi-sterile to a greater or lesser extent, their corrected fertility ranging from 5.0% to 57.0%. The mean fertility of 23.0% differs from that given for the dose-response curve in which bugs given 6 K rad had a mean fertility of 44.8%, but the latter figure was based on only 4 pairs of bugs.

Table 4.3. Fertility data for the parental generation of 40 virgin male *B. prolixus* irradiated with 6 K rad and mated with normal virgin females. The data for eggs laid after 4 blood-meals have been pooled. Control fertility = 91.88%

Pair number	Number of eggs laid	Number of eggs hatched	Corrected % fertility
1	62	17	29.84
2	85	25	36.84
3	47	11	25.47
4	29	7	26.27
5	58	16	33.78
6	71	7	10.73
7	0	0	0
8			
9	80	14	19.04
10	61	11	19.63
11	56	8	7.78
12			
13	59	14	25.83
14	38	7	20.05
15			
16	22	2	9.89
17			
18	77	11	19.79
19	51	6	12.80
20	14	0	0
21			
22	Male died		
23	Male died		
24	Female died		
25	19	8	22.91
26	14	3	15.55
27			
28	38	4	11.46
29	62	23	57.93
30	43	14	35.44
31	37	8	23.53
32	32	6	20.41
33	60	18	25.40
34			
35	66	12	19.79
36	45	7	16.93
37	49	3	15.55
38	101	20	26.94
39	31	6	20.00
40			
	35	14	43.54
<u>TOTAL</u>	<u>1423</u>	<u>313</u>	<u>29.88</u>

eggs laid after each of the 4 parental blood-meals were kept separately, and, on hatching, the larvae were reared to F_1 adults as 4 separate batches derived from egg batches 1, 2, 3 or 4. All of the offspring from these 28 partially fertile pairings were fed but not all of them were successfully reared to adults. Of the 31 first instar larvae which hatched, only 147 were reared to adults (43.1%). The majority of the 147 bugs which were fed reared successfully died during the early stages of development as first or second instar larvae. Of the 147 bugs reared to adults 86 were males and 61 females, and a χ^2 test showed that there was no significant departure from an expected 1:1 sex ratio in these F_1 bugs ($\chi^2 = 3.4$, $P = N.S.$).

Only 86 of the 147 F_1 adults were used in further matings. Selection of these 86 bugs was made to include bugs from families with few surviving progeny and to limit the number of bugs used from large F_1 families in which survival to F_1 adults had been good; thus the choice of parents was not random. These 86 selected adults were each mated with normal males and given 3 blood-meals at 14 day intervals. The hatchability data of the eggs laid by these 86 F_1 pairs were pooled for all eggs laid by each pair over 3 blood-meals (Tables 4.4 - 4.7 - each of these 4 Tables relating to crosses of normal adult $\times F_1$ adults from the egg batches 1 to 4 laid by the parental female \times irradiated parent male cross).

The results show that 2 of the 86 $F_1 \times$ normal matings were fully fertile (No. 2₁ from the first egg batch and No. 11 from the third egg batch) and that both of these were $F_1 \times$ normal crosses. Both three of the $F_1 \times$ normal crosses were completely sterile and if we assume sterility, the semi-sterile crosses showing a wide range of fertility from 1/250 from 11₁ egg batch No. 41 to 1/1000 from 2₂ egg batch No. 25.

Table 4.4. Fertility data for 14 crosses of F_1 X normal R. prolixus, the F_1 adults being derived from egg batch 1 of normal female parent X irradiated male parent. Data pooled for eggs laid after 3 blood-meals by F_1 females.
Control fertility = 91.88%

Bug number P	F_1	F_1 sex	Number of eggs laid	Number of eggs hatched	Corrected % fertility
1	1	♂	31	0	0
	2	♀	94	51	59.05
2	1	♀	132	114	94.00
	2	♀	120	0	0
4	1	♂	85	13	16.65
	2	♀	143	5	3.88
	3	♂	86	3	3.60
9	1	♀	124	3	2.63
19	1	♂	74	24	35.30
29	1	♂	79	25	34.44
	3	♂	63	2	3.20
30	1	♂	76	45	54.44
	2	♂	61	5	8.92
34	1	♀	62	2	3.51
Total			1230	292	25.94

Table 1-1. Fertility data for *Phormica* of F_1 X normal *S. mellonae*, the F_1 adults being derived from egg batch 2 of normal female parent X irradiated male parent. Data pooled for eggs laid after 3 blood-meals by F_1 females.
Control fertility = 91.88%

Seg number F	F_1	F_1 sex	Number of eggs laid	Number of eggs hatched	Corrected % fertility
3	3	0	48	0	2.27
	4	0	87	0	1.33
	5	0	39	0	0
	6	0	81	0	0
7	7	0	59	11	20.29
	8	0	72	43	71.05
	9	0	87	0	0
	10	0	33	0	0
11	11	0	165	0	0
	12	0	27	0	0
	13	0	100	0	9.80
14	14	0	27	0	0
	15	0	95	0	4.58
	16	0	37	0	0
	17	0	44	23	28.91
18	18	0	61	0	0
	19	0	38	0	0
	20	0	66	1	1.65
	21	0	71	1	1.53
22	22	0	25	0	0
23	23	0	4	0	0
24	24	0	51	0	6.40
25	25	0	56	1	1.94
26	26	0	110	0	0
27	27	0	12	0	0
28	28	0	66	1	1.55
29	29	0	22	0	0
30	30	0	67	0	45.48
31	31	0	30	0	0
32	32	0	70	1	1.55
33	33	0	135	0	55.63
34	34	0	117	0	3.72
35	35	0	113	11	11.51
36	36	0	45	3	7.62
37	37	0	115	0	35.96
38	38	0	70	1	1.55
39	39	0	50	0	0
40	40	0	40	0	0
Total			2024	207	11.19

Table 4.6. Fertility Data for 20 females of F_1 Normal C. quinquefasciatus. The F_1 adults being derived from egg batch 3 of normal female parent X irradiated male parent. Data pooled for eggs laid after 3 blood-meals by F_1 females. Control fertility = 91.88%.

Egg number # - F_1	F_1 sex	Number of eggs laid	Number of eggs hatched	Corrected % fertility
2	♀	179	0	0
5	♀	147	3	2.04
	♀	64	0	0
10	♀	120	0	0
13	♀	49	0	0
	♀	138	102	73.91
18	♀	71	0	0
28	♀	71	0	0
	♀	120	0	0
	♀	81	0	0
	♀	94	0	0
	♀	134	0	0
	♀	37	0	0
	♀	32	0	0
34	♀	60	0	0
	♀	69	0	0
36	♀	47	0	0
	♀	12	0	0
37	♀	45	0	0
	♀	104	1	0.96
Total		1784	103	5.77

Table 2.7. Fertility data for 13 crosses of F_1 X normal *A. aroclinus*, the F_1 adults being derived from egg batch 4 of normal female parent X irradiated male parent. Data pooled for eggs laid after 3 clonings of F_1 females. Control fertility = 91.88%.

Bug number P	F_1 sex	Number of eggs laid	Number of eggs hatched	Expected % fertility
2	1	♂	46	0
	2	♂	70	0
3	1	♀	139	0
	2	♀	26	0
13	1	♂	91	13.55
	3	♀	142	0
25	1	♀	102	0
29	2	♀	9	0
37	1	♂	40	0
	3	♀	116	0
40	1	♂	51	0
	2	♀	89	1.22
	3	♂	86	26.58
Total		1001	75	8.78

Table 4.8. F_1 X normal *R. prolixus* fertility data pooled from 4 parental egg batches (see Tables 4.4 - 4.7) representing 22 parental crosses. Control fertility = 91.88%.

Parental family No.	Number of F_1 pairs in sibship	Number of eggs laid	Number of eggs hatched	Corrected % fertility
1	8	375	53	15.38
2	8	172	172	23.46
3	8	330	0	0
4	8	36	0	7.40
5	8	211	0	1.55
6	8	124	10	8.78
7	8	256	25	10.63
8	8	120	0	0
9	8	0	0	35.76
10	8	236	2	0.92
11	8	71	0	0
12	8	99	24	26.38
13	8	102	0	0
14	8	0	24	3.96
15	8	151	27	19.46
16	8	137	50	39.72
17	8	166	1	0.66
18	8	0	0	0.93
19	8	59	0	0
20	8	862	106	13.38
21	8	343	55	17.45
22	8	326	35	11.68
Total for 22 sibships		6520	757	12.64

The data from these four groups presented in Tables 4.2 + 4.3 have been pooled and are presented in Table 4.4 as data from 14 families.

The results show that the mean corrected fertility for the 86 F_1 X normal crosses was $12.6\% \pm 0.04$. Five of the 22 F_1 sibships were completely sterile and the mean fertility of the other 17 sibships ranged from 1.0% to 75.7%. These figures may be compared with the parental fertility of 23.0% range 1.0% to 41.0% which indicated that the F_1 progeny were much less fertile than their irradiated parents. However, a straight-forward comparison of these F_1 mean values is not meaningful since the F_1 ratings were derived from several different parents, but 27 F_1 crosses from 8 crosses of the range of fertilities for the 2 generations that the F_1 families were less fertile overall. A family by family comparison of the fertility data from Tables 4.3 and 4.4 shows that in only 3 of the 10 families was the F_1 fertility greater than that of the corresponding parental family. The 3 exceptions, families 12, 13 and 14 had increased F_1 fertility but χ^2 tests of the data showed that these increases were not significant over parental fertility ($\chi^2 = 0.98, 1.91$ and 3.41 respectively, $P > N.S.$).

A comparison of the fertility of F_1 ratings established from each of the 4 parental egg batches revealed that the fertility of these 4 groups differed. The mean fertility of the 4 groups listed in Table 4.2 + 4.3 were 25.0%, 22.0%, 11.1% and 5.1% respectively, and the F_1 bugs from the first parental egg batch were more fertile than those of the later batches. A simple statistical comparison of these 4 proportions would not be meaningful since not all F_1 sibships were represented in each of the 4 parental egg batches. A comparison was therefore made between F_1 families within sibships for the 4 batches

The data from these four groups presented in Tables 4.2 - 4.7 have been pooled and are presented in Table 4.8 as data from 22 families.

The results show that the mean corrected fertility for the 22 F_1 4 normal crosses was $10.8 \pm 5.4\%$. Five of the 22 F_1 sibships were completely sterile and the mean fertility of the other 17 sibships ranged from 0.0% to 25.7%. These figures are in comparison with the parental fertility of 13.8% (range 0.0% to 27.0%) which indicated that the F_1 progeny were much less fertile than their irradiated parents. However, a straight-forward comparison of these 2 mean values is not meaningful since the F_1 matings were derived from several different parents, but it is clear that a mean of the range of fertilities for the 2 generations that the F_1 families were less fertile overall. A family by family comparison of the fertilities data from Tables 4.3 and 4.8 shows that in only 3 of the 22 families was the F_1 fertility greater than that of the corresponding parental family. The 3 exceptions, families 13, 19 and 20 had increased F_1 fertilities but χ^2 tests of the data showed that these differences were not significant over parental fertility ($\chi^2 = 0.05, 0.46$ and 1.11, respectively, $P = N.S.$).

A comparison of the fertility of F_1 matings established from each of the 4 parental egg batches revealed that the fertility of these 4 groups differed. The mean fertilities of the 4 groups shown in Tables 4.4 - 4.7 were 25.8%, 11.5%, 5.1% and 2.0% respectively, and the F_1 bugs from the first parental egg hatch were more fertile than those of the later batches. A simple statistical comparison of these 4 proportions would not be meaningful since not all F_1 sibships were represented in each of the 4 parental egg batches. A comparison was therefore made between F_1 families within sibships for the 4 batches

Table 4.9. χ^2 tests for differences in fertility between 62 F_1 families of *B. prolimus* grouped into 11 sibships and produced from 4 egg batches laid by the parental generation.

F ₁ sibship number	Parental egg batch								df	X ²	P
	1		2		3		4				
	L	H	L	H	L	H	L	H			
1	125	51	250	2					1	75.8	<0.001***
2	252	114	251	58	179	0	116	0	3	108.9	<0.001***
4	314	21	127	9					1	0.02	N.S.
9	124	3	132	22					1	12.2	<0.001***
14			236	2	187	125			1	116.9	<0.001***
19	74	24	25	0					1	7.6	0.01-0.001**
28			91	3	569	21			1	0.02	N.S.
29	142	27					9	0	1	1.69	N.S.
34	62	2	12	0	159	0			2	5.4	0.1-0.05
37			507	105	199	1	156	0	2	65.7	<0.001***
40			90	0			226	22	1	8.5	<0.001***
Total	1093	242	1886	201	1293	147	507	22			

L = No. of eggs laid

H = No. of eggs hatched

df = Degrees of freedom

P = Probability

and the results of these analyses are shown in Table 4.9. Only those 11 sibships which were each represented for more than one parental batch could be compared and the results show that there were significant differences in fertility between families for 7 of these 11 sibships. These analyses do not reveal whether there was any uniformity or trend in these differences between egg batches, therefore a test for linear trend in proportions was carried out on the column totals given in Table 4.9, the results of which are shown in Table 4.10.

Table 4.10. χ^2 test for linear trend in fertility for 11 F_2 families from 4 parental egg batches.

	Parental egg batch				χ^2	d.f.	P
	1	2	3	4			
No. eggs laid	1093	1886	1293	547	-0.04	18	<0.001***
No. eggs hatched	42	201	189	10	-0.01	18	<0.001***

b = slope

The results show that there was a highly significant linear trend in proportions downwards from egg batch No. 1 to egg batch No. 4 ($\chi^2 = -0.04$, $P < 0.001$), and it may be concluded that F_1 eggs from the 4 parental crosses had significantly different fertilities, with a trend towards reduced fertility in the later batches, reflecting differences due to birth order of these F_1 males. Fertilization of these 40 F_1 female midges were completely sterile compared with only 10 of the 40 parental midges and the proportions of sterile, semi-sterile and fully fertile F_2 females

specimens have been maintained for each of the 4 batches. The results of which are shown in Table 4.11.

Table 4.11. Proportions of sterile, semi-sterile and fully fertile bugs for 86 F_1 X normal matings from 4 parental egg batches.

Parental egg batch	No. of F_1 pairs	Sterile		Semi-sterile		Fully fertile	
		No.	%	No.	%	No.	%
1	14	2	14.3	11	78.6	1	7.1
2	39	18	46.2	21	53.9	0	0
3	20	13	65.0	6	30.0	1	5.0
4	13	10	76.9	3	23.1	0	0
Total	86	43	50.0 ± 0.8	41	46.7 ± 0.7	2	2.3 2.2

The results show that 47% (41) of the 86 matings were fully fertile, 50% (43) were completely sterile and 2% (2) were semi-sterile. Furthermore, it is clear that the proportions of completely sterile bugs increased with birth order from 14.3% in egg batch 1 to 76.9% in egg batch 4, while the proportions of semi-sterile bugs decreased from 78.6% in egg batch 1 to 23.1% in egg batch 4. These data indicate a strong correlation between birth order of F_1 bugs and the proportions carrying dominant lethal mutations.

An interesting comparison was made between F_1 ♂ X normal ♀ and F_1 ♀ X normal ♂ fertilities. In such a comparison it would not be valid simply to contrast the mean percentage egg hatch of the 2 groups since the analysis must allow for family relationships within groups, i.e. ♂ and ♀ F_1 parents of the same sibship must be compared with each other. The method adopted for this comparison was a weighted logistic analysis, the details of which are given in Cox (1973) Chapter 3. The data for this analysis were shown in Table 4a,b and the results of the calculations are shown in Table 4c,d.

Table 4c,d. Weighted logistic analysis of F_1 ♂ X normal ♀ and F_1 ♀ X normal ♂ fertility data. Only fertile or semi-fertile crosses included.

Cross	No. of pairs	No. of eggs laid	No. of eggs hatched	% fertility	Weight	Estimate	Z
F_1 ♂ X normal ♀	24	1787	277	15.5	-220.884	28.1001	3.63 <0.001***
F_1 ♀ X normal ♂	19	1977	461	23.6	-200.477	26.2524	

Z = Standard normal deviate

The results show that in a comparison of 43 fertile or semi-fertile crosses, F_1 female lines were significantly more fertile to some 51 egg fertilities than F_1 male lines were able to achieve. However, since 10 of the 19 ♀ lines which included pairs from the F_1 ♂ X normal female matings,

only 153 (55.2%) came through successfully to adults as against 320 (70.7%) reaching adult of the 467 larvae hatching from the 19 P_1 \bar{W} X normal male matings.

The sex of these ' P_2 ' bugs was tested for departures from the 1:1 expected ratio and the results showed that the proportions of male to female was almost exactly 1:1; viz., 153 bugs raised from P_1 \bar{W} X normal \bar{Q} 79 were male and 74 were female, and, of the 80 bugs raised from P_1 \bar{W} X normal \bar{Q} 162 were male and 168 were female.

Twenty-eight of these ' P_2 ' bugs were further crossed with normal males, those 28 being those from among the 31 P_1 crosses with the lowest fertilities. These ' P_2 ' pairings were each given 3 blood-meals and the eggs produced were scored for hatchability (Table 4.14). The results show that the mean fertility of young of selected ' P_2 ' 28 crosses was 45.8%, with family fertilities ranging from 1.0% to 100.0%. The fertility of 20 of the ' P_2 ' 28 crosses was superior to that of their respective parents' fertilities except for families 30, 31, and 28, which had slightly reduced fertilities in comparison with their parents.

Surprisingly, 6 of the 28 ' P_2 ' crosses were fully fertile even though all crosses were deliberately selected from the P_1 families with the lowest fertilities. For comparative purposes the mean fertilities and ranges in fertility for all families raised from parental, P_1 and ' P_2 ' crosses are shown in Table 4.14.

Table 4.13. Fertility of 28 'P₂' X normal crosses, data pooled for all eggs laid after 3 blood-meals. Parental and grand-parental fertilities are included for comparison.

Family No.			Sex	Eggs		Corrected % fertility			Parental egg batch
P	F ₁	F ₂		Laid	Hatched	F ₂	F ₁	P	
4	2	1	♂	25	19	87.72	1.81	26.27	1
			♂	54	22	36.78	3.81	26.27	
			♂	54	3	6.05	3.80	26.27	
			♂	39	16	20.67	3.83	26.27	
		1	♂	67	15	41.86	3.46	35.44	
			♂		5	8.77	8.92	3.53	
6	2	1	♂	77	8	11.30	4.58	10.73	2
			♂	59	57	100.00	6.40	26.94	
		2	♂	87	6	7.50	6.40	26.94	
		2	♂	102	100	100.00	6.40	26.94	
		4	♂		died				
		5	♂	80	3	4.08	3.72	26.94	
5	1	1	♂	47	18	41.68	5.22	11.78	3
	2	1	♂	59	5	9.22	8.16	57.93	
			♂	13	13	23.98	8.16	57.93	
		1	♂	70	17	26.43	3.47	57.93	
			♂	39	1	2.79	3.47	57.93	
			♂	22	4	19.79	3.47	57.93	
		1	♂	56	6	11.66	4.87	57.93	
			♂	48	47	100.00	4.87	57.93	
		3	♂	54	0	0	4.87	57.93	
		4	♂		died				
37	2	1	♂	100	93	100.00	1.05	26.94	
11	1	1	♂	63	61	100.00	15.55	25.80	4
			♂	36	8	24.19	15.55	25.80	
	3	2	♂	33	9	29.68	26.85	43.54	
			♂	69	62	97.80	26.85	43.54	
		4	♂	49	47	100.00	26.85	43.54	
Total				1609	660	44.64	5.44	28.58	

Table 4.4. Corrected fertility for 3 generations of families of H. prolixus to compare fertility of all families with fertility of selected families.

Generation	All families		Selected families	
	Mean fertility %	Range %	Mean fertility %	Range %
P	23.9	7.9 - 57.9	28.6	10.7 - 57.9
F ₁	12.6	1.1 - 28.6	5.4	1.1 - 20.9
'F ₂ '	44.6	2.8 - 100	44.6	1.8 - 100.0

The results show that the 16 F₂ families selected for breeding had a mean fertility of 44.6 (range 1.8 - 100.0) which was significantly higher than the overall mean for F₂ families of 12.6 (range 1.1 - 28.6). These data show not only that following irradiation with 6 K and 7-rays H. prolixus males had a much reduced fertility compared with control matings, but also that the fertility of the F₂ progeny was raised with respect to the mean of the total F₂ population. Furthermore, when 16 F₂ bugs specifically selected from under F₁ females of the fertility were crossed with normal males, the fertility of this 'F₂' generation is seen to have returned substantially to control.

Only a few 'F₃' bugs were reared to adults and these all came from 4 'F₂' families descended from the three original F₂ bugs. These bugs were not mated and therefore no fertility data were produced for this generation.

Cytogenetic studies

It is now appropriate to consider the cytogenetic findings with

relation to these fertility data, and those were derived from parentage bugs of the above experiments, after male parents had been withdrawn from their mating period. Prole 441 (diploid) metaphase I nucleus from a normal male spermatocyte of *M. jelskii*, the diploid number of which is $2n = 20$ autosomes + XY. The karyotype of *M. jelskii* as first described by Uehime (1966) to consist of 3 longer pairs of autosomes and 7 shorter pairs, although it was not possible from that investigation to distinguish individual bivalents, in meiotic preparations, only the sex univalents being readily identifiable. The 10 pairs of autosomes are usually associated by a single centromere to form a bivalent.

Lacto-acetic-orcein squash preparations were made from the fixed testes of F_1 males, their sons (F_2) and grandsons (F_3). Forty-nine of the 51 F_1 males were examined cytogenetically after mating with normal females; the other 2 males were like parents because of sterility. Eighty-nine F_2 males were examined cytogenetically, of these F_2 48 + normal + autosomes and 10 from F_2 10 + normal + autosomes. Only 11 F_3 males were used for cytogenetic examination. In scanning the testis squashes any departures from the normal 10 bivalents + XY configurations at metaphase I were noted and photographs taken for compiling cytogenetic data.

All 49 F_1 males examined revealed chromosomal abnormalities in their spermatocytes so that in every male at least one interchromosomal aberration was present. While many males showed 2 abnormalities and some had 3. In addition, normal cells showed normal pairing chromosomes fragments, presumably detached from highly fragmented bivalents. The cytogenetic results obtained by examination of F_1 male testes have been summarized and are shown in Tables 2, 3 & 4. Together with fertility data. It must be emphasized that although patterns of

Table 2.15. Chromosomal associations at metaphase I in F_2 male *P. variator* from disassociated male parents

X normal female parent: - first egg batch. Except where indicated, multivalents associated as chains.

F_2 male number	Metaphase I associations	Abnormalities	% fertility F_2	Parental fertility
1,1	8 II + 1 III + I	1 Translocation	0	29.8
4,1	8 II + 1 III + I	1 "	16.6	26.3
4,1	6 II + 2 III + 2 I	2 "	3.7	26.3
19,1	6 II + 2 III + 2 I + F	2 "	35.3	14.8
29,1	8 II + 1 III + I	1 "	34.4	35.4
29,1	8 II + 1 III + I	1 "	3.5	35.4
30,1	8 II + 1 III + I	1 "	64.4	23.5
30,2	8 II + 1 III + I	1 "	8.9	21.5

F = Fragments

Table 4.14. Chromosomal associations at metaphase I of F_2 male *S. similis* from irradiated male parents

X normal female parents - second egg batch. All multivalents associated as chains, except where stated.

F_1 male number	Metaphase I associations	Abnormalities	% fertility F_2	Parental fertility
1,1	6 II + 1 IV + 1 III + 1 I + 1 X	2 Translocations	2.3	29.8
2,6	8 II + 1 III + 1 I	"	0	36.8
6,1	8 II + 1 III + 1 I	"	20.6	10.7
7,1	4 II + 3 III + 3 I	"	0	19.0
8,3	8 II + 1 III + 1 I + 1 X	"	7.1	19.2
11,1	6 II + 1 IV + 1 III + 1 I	"	0	21.0
14,2	6 II + 2 III + 2 I + 1 X	"	0	20.2
20,1	4 II + 1 IV + 2 III + 2 I	"	0	57.9
28,2	6 II + 2 III + 2 I	"	6.4	57.9
32,1	8 II + 1 III + 1 I	"	1.0	25.4
34,2	8 II + 1 III + 1 I	"	0	19.8
35,1	8 II + 1 III + 1 I	"	4.9	26.9
37,2	6 II + 2 III + 2 I	"	0	26.9
37,3	6 II + 1 IV + 2 III + 1 III + 1 I	"	45.5	26.9
37,4	6 II + 2 III + 2 I	"	0	26.9
37,5	6 II + 2 III + 2 I	"	1.6	26.9
38,2	8 II + 1 IV	"	7.2	28.1
40,1	6 II + 2 III + 2 I	"	0	43.5
40,2	8 II + 1 III + 1 I	"	0	43.5

Table 4.17. Chromosomal associations at metaphase I in F_1 male *S. pomonella* from irradiated male parents

X normal female parents - third egg batch. All multivalents associated as chains except where stated.

F_1 male number	Metaphase I associations	Abnormalities	% fertility F_1	Parental fertility
5,4	6 II + 2 III + I	2 Translocations	0	11.8
10,1	8 II + 1 III + I	1 "	0	19.6
13,1	6 II + 2 III + 2 I	2 "	0	25.8
18,1	6 III + 2 III + 2 I	2 "	0	19.8
28,1	8 II + 1 III + I	1 "	4.6	25.8
28,2	8 II + 1 III + I	1 "	8.2	57.9
28,3	8 II + 1 IV	1 "	0	57.9
28,4	6 II + 2 III + 2 I	2 "	3.5	57.9
28,5	6 III + 2 III + 2 I	2 "	4.9	57.9
28,6	8 III + 1 IV	1 "	0	57.9
28,7	8 III + 1 III + I	1 "	0	57.9
34,2	8 II + 1 IV	1 "	0	19.8
34,3	8 III + 1 III + I	1 "	0	19.8
36,2	6 II + 1 IV + 1 III + I	2 "	0	15.5
37,1	8 III + 1 III + I	1 "	0	26.9
37,2	8 II + 1 III + I	1 "	1.3	26.9

Table 4.18. Chromosomal associations at metaphase I in testis squashes of F_1 male *S. prolixus* from irradiated male parents X normal female parents - fourth egg batch.

F_1 male number	Metaphase I associations	Abnormalities	fertility F_1	Parental fertility
2	8 II + 1 III + I	1 Translocation		36.8
13	4 II + 1 III + I + 2 IV	3 "		36.8
13	8 II + 1 III + I	"	15.5	25.0
17	6 II + 2 III + 2 I	2 "	0	26.0
40	4 II + 3 IV	3 "	0	43.0
40	8 II + 1 III + I	"	6.5	43.0

Table 4.9. Metaphase I associations and translocation frequencies for 49 *F₁* males from 4 parental egg batches (*R. prolixus*).

No. of translocations	Metaphase I associations	Parental egg batch No.				Total	%
		1	2	3	4		
1	8 II + 1 III chain + 1	6	7	7	3	23	46.9
	8 II + 1 IV chain		1	3		4	8.2
	8 II + 1 IV ring						
	Sub total	6	8	10	3	27	55.1
2	6 II + 2 III chains + 2 I	2		5	1	14	28.6
	6 II + 1 IV chain + 1 III chain + I		1	1		3	6.1
	6 II + 1 IV ring + 1 III chain + I		1			1	2.0
	Sub total			6	1	10	16.7
3	4 II + 3 III chains + 3 I		1			1	2.0
	4 II + 1 IV chain + 2 III chains + 2 I		1			1	2.0
	4 II + 3 IV chains				1	1	2.0
	4 II + 2 IV chains + 1 III chain + 1				1	1	2.0
	Sub total	0	2	0	2	4	8.2
	Total for 4 egg batches	8	19	16	6	49	

metaphase I chromosome associations often showed some degree of variation between cells within individual testis squashes; no attempt has been made to quantify this variation; the associations presented in the Tables of results represent either the only pattern seen or else the most common association within a particular testis. The numbers of translocations per individual shown in Tables 4.15- 4.18 were inferred from the patterns of association seen at metaphase I. When inter-batch data were used to compare the translocation frequencies of the F_1 males from the 4 parental egg batches. The translocation frequencies are shown in Table 4.19. The results show that of the 49 males examined, 27 (55.1%) showed a single translocation, 18 (36.7%) showed evidence of 2 translocations and 4 bugs (8.2%) showed evidence of having 3 translocations. Analysis of the distribution of translocations within parental egg batch did not reveal any statistically significant differences, probably because of the small numbers in each group, but it can be noted by inspection, that the majority of males from the first egg batch had only a single translocation while in later batches 2 and 3, 2 translocations seemed more frequent.

The most striking feature of these analyses of multivalent associations is that of the 49 F_1 males examined, only one bug showed a ring multiple association (bug No. 37, from egg batch No. 2), the rest all having multivalents associated at metaphase I as chains of III plus quadrivalents, or chains of III and IV. The production of these chain multiples may be explained as represented in Fig. 4.1 = 4.4. It has already been explained in the introduction, how the chromosomes of an F_1 male, heterozygous for a single interchange must form a Zygotene cross at meiosis to ensure regular pairing (Fig. 4.1), but the pattern of association at metaphase I will depend on numbers and

positions of chiasmata formed within the cross, and the numbers of chiasmata formed will in turn be dictated by the morphology of the chromosomes involved in the associations. If chiasmata are formed in all 4 pairing segments of the zygotene cross, a ring of 4 chromosomes will be produced in metaphase I as illustrated in Fig. 4.11a. It must be noted that, however, as in the interstitial aspect (i.e., the section of a multiple lying between the centromere and point of exchange in a monocentric multiple) in multiples formed by holokinetically organisms, so that the special associations produced by interstitial crossing-over (Lewis and John, 1963) need not be considered here.

The genetic consequences of ring formation are illustrated in Fig. 4.11b. The standard mode of nomenclature used for monokinetically organisms has been adopted here, although the terms "alternant" and "adjacent", which they define the chromosome positions, cannot strictly be applied to holokinetically species. Although no centromeres are illustrated in these line drawings, it is still clear that only if "alternant" ends of orientation can lead to the production of genetically balanced gametes and that this involves a twisting of the multiple at metaphase I.

If, however, one of the arms of a zygotene cross is short, then chiasma formation may fail in that arm, resulting in the production of a chain of IV at metaphase I (Fig. 4.11c). Other than a ring multivalent. Similarly, if there are two consecutive short arms in the zygotene cross, there may be a failure in the formation of two chiasmata, resulting in a chain of III plus a univalent (Fig. 4.11d). It also follows that if 2 short arms of the zygotene cross alternate with 2 long arms, then chiasma formation in the long arms alone could result in the production of 2 bivalents at metaphase I. The arrangement of chromosomes within a chain will obviously depend on which arms

of the Xygotone cross are chiasmata. Thus, for a single interchange there are 4 possible arrangements of the chromosomes in a chain multiple and these are shown in Fig. 4.3(1) for both a chain of IV and a chain of III + I. The genetic consequences of chain formation will, however, depend on the orientation of the multiple in relation to the spindle at metaphase I. The possible modes of orientation of chains of IV and III + I are illustrated in Fig. 4.4. Lewis and Jones (1963) have shown that it is possible for a chain of IV to orientate in 4 ways: linear, convergent, indifferent and parallel, while a trivalent chain of III + I can only orientate in 3 of these ways, the parallel mode being impossible. When disjunction of these chain multiples occurs, only the convergent orientation type leads to the production of balanced gametes (Lewis and Jones, loc. cit.).

It may be seen from the summary of results presented in Table 4.1 that the most frequent association found in *P. pallidus* ♂ males with a single interchange was $2\text{ IV} + 1\text{ III chain} + 1\text{ X}$. Since, for reasons, the comparative purposes, a normal metaphase I cell from adult male *P. pallidus* with 10 bivalents gave 10. Table 4.1 shows a metaphase spermatocyte from P_1 ♂ No. 4₁ (egg batch No. 1) with a single interchange associated as a chain of III + I, the chain being linearly orientated.

Four P_1 males were found with chain of IV associations of a single interchange, one of which is illustrated in Table 4.1 which comes from P_1 ♂ No. 38₁ (egg batch No. 2) with chain of IV linearly orientated. Not a single ring of IV association was found in any of the P_1 males with a single translocation.

The results given in Table 4.1 show that 10.7% of the P_1 males examined showed metaphase configurations which indicated that their chromosomes were involved in 2 interchanges. Plate 4.4 shows a metaphase I spermatocyte from P_1 bug No. 19₁ (egg batch No. 1) associated

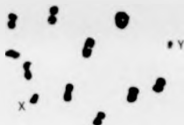


Plate 4-1. Normal metaphase I spermatocyte from unfertilized adult male

Hy. mull. (1949) Slide 15 (Kalamata 1949) X 2000.



Plate 4-2. Metaphase I spermatocyte from Hy. mull. (1949) Slide 15 (Kalamata 1949)

showing single translocation associated as $8 \text{ II} + 1 \text{ III} + \text{I}$,
one translocation between chromosomes. (Kalamata 1949) X 2000.



Plate 4.3. Metaphase I spermatocyte from F_1 male (No. 382) *R. prolixus* showing single translocation associated as bivalent. One chain of bivalent orientated. Fertility = 7.2%. X 2500.



Plate 4.4. Metaphase I spermatocyte from F_1 male (No. 193) *R. prolixus* showing two translocations associated as bivalents. One chain orientated linearly (I) the other orientated circularly (II). Fertility = 34.4%. X 2500.

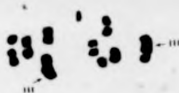


Plate 4.5. Metaphase I spermatocyte from F_1 male (No. 37₄) R. prolixus showing two translocation both associated as linear chains of III + I. Fertility = 0%. X 2500.



Plate 4.6. Metaphase I spermatocyte from F_1 male (No. 14₁) R. prolixus showing two translocations associated as (i) chain of IV and (ii) chain of III + I. The chain of IV shows parallel orientation and the chain of III indifferent orientation. Fertility = 0%. X 2500.

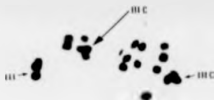


Plate 4, 9. Metaphase I spermatocyte from 17th male (No. 38, 143) 8. 11. 1955.
 Showing three translocations associated with chains of III + X, one being linearly orientated, the others showing univalent associations. 8 PMN.



Plate 4, 10. Metaphase I spermatocyte from 17th male (No. 62, 11) 8. 11. 1955.
 Showing two translocations, one being a chain of IV in parallel orientation, the other a chain of III + I in parallel orientation. Fertility 11.30. 8 PMN.



Figure 4.11. Chromosome 1 squamous cell carcinoma (T₁) male fib. (S₁, S₂) showing
 showing - single chromosomal fragment. Fertility - 0%.
 X 2500.



Figure 4.12. Chromosome 1 squamous cell carcinoma (T₁) male fib. (S₁, S₂) showing
 showing two chromosomal fragments. Fertility - 41.7%. X 2500.

as $6 \text{ II} + 2 \text{ III} + 1 \text{ I}$, the chains of III showing different forms of orientation, one being linear the other bidirectional. Plate 4.7 shows a cell from F_1 male 37_4 (egg batch No. 2) with 2 translocations in the form $6 \text{ II} + 2 \text{ III} + 1 \text{ I}$ but in this case both chains of III are linearly orientated.

Three F_1 males showed chain of IV associations together with a chain of III plus a univalent in the same cell. This type of configuration involving 2 translocations is illustrated by Plate 4.6 which shows a metaphase I cell from F_1 male No. 14_1 (egg batch No. 2) in the association $6 \text{ II} + 1 \text{ IV} + 1 \text{ III} + 1 \text{ I}$, the chain of IV showing parallel orientation and the chain of III showing bidirectional orientation.

Only one cell was found showing a multivalent associated with a ring, and this was in bug No. 37_3 (egg batch No. 2) which had 1 translocation. Plate 4.7 shows a metaphase I spermatocyte from F_1 male 37_3 with a ring of IV and a linear chain of III plus a univalent, the ring showing the twisted configuration necessary for alternate orientation. The formation of such ring multivalent association, although not observed here in the present experiment, is important since it demonstrates that ring formation is not precluded in *D. melanogaster*.

Four F_1 males were found to show evidence of 3 translocations, all of which showed chains with linear orientation. Plate 4.8 shows a metaphase I cell from F_1 bug No. 5_2 (egg batch No. 1) with the association $4 \text{ II} + 3 \text{ III} + 3 \text{ I}$, the chains all linearly orientated.

As the photomicrographs develop, the mode of association of multivalents at metaphase I and also their orientation in the nucleus becomes variable. This variation occurring between cells within tissues, the results show that the most frequent association in these F_1 males was of the form $6 \text{ II} + 1 \text{ III} + 1 \text{ I}$ suggesting that most of the viable

translocations produced in the parental males by irradiation resulted in the formation of synaptonemes crossed with 2 short arms which could result in the failure of chiasma formation in these arms and the subsequent chain of III plus univalent associations. The photomicrographs also show that the majority of the chains were orientated linearly on the metaphase spindle as shown in Plate 4.4.

The cytogenetic results from ' P_2 ' male testis squash preparations are presented in Tables 4.20 - 4.25 and Table 4.26 contains a summary of these results.

The summary of the ' P_2 ' male cytogenetic data is shown in Table 4.26 and of the 34 males examined, 34 (100%) had no detectable chromosomal abnormalities, which is in sharp contrast to the ' P_1 ' males which all showed some degree of abnormality. Fifty-five of the 80 ' P_1 ' males (68.75%) however, were evidence of chromosomal abnormalities, viz. 10 (18.2%) with a single translocation, 8 (14.5%) with 2 translocations, and one with 3 translocations. These figures are more different from the ' P_2 ' males, 36.7% of which had 2 translocations and 34.7% had 3 translocations. (Table 4.2 shows a metaphase 2 karyotype from the only ' P_1 ' male with 3 translocations (no. 90, 36.7%) associated as 4 (I - 3, II + 3, I), one chain showing linear orientation but the other 2 showing convergent orientation. The observation of nuclei showing convergent orientation is important for, as already stressed, only this type of chain orientation may lead to the production of balanced gametes (see Fig. 4.4) which are essential for the maintenance of a translocation in a population.

Twenty-eight of the 34 ' P_2 ' males with a single translocation were found to be associated in the form of a chain of III plus a univalent while 6 were associated as a chain of IV. The modes of association of these chains in ' P_2 ' males again showed variation both between and within

Table 4.20. Chromosomal associations at metaphase I in testis squashes of 'F₂' male B. pumilio from F₁ male (first egg batch) X normal female. All multivalents associated as chains except where stated.

Bug number F ₁ F ₂	Metaphase I associations	Abnormalities	% fertility F ₂	F ₁ parental fertility %
4 ₁ 2	All cells normal	None		
3	8 II + 1 III + I/8 II + 1 IV	1/2 Translocations		
4	8 II + 1 IV	1 "		
4 ₃ 1	8 II + 1 III + I	1 "	6.0	3.7
19 ₁ 1	1 Fragment	1 Fragment		
2	8 II + 1 III + I	1 Translocation		
29 ₁ 4	All cells normal	None		
5	8 II + 1 III + I	1 Translocation		
6	All cells normal	None		
8	All cells normal	None		
30 ₁ 1	All cells normal	None		
12	8 II + 1 III + I	1 Translocation		
14	All cells normal	None		
16	8 II + 1 III + I	1 Translocation		
17	All cells normal	None		
30 ₂ 2	8 II + 1 III + I	1 Translocation	6.8	6.9
30 ₃ 1	All cells normal	None	41.9	3.5

Table 4.21. Chromosomal associations at metaphase I in testis squashes of 'F₂' male B. prolixus from F₁ female (first egg batch) X normal male. All multivalents associated as chains except where stated.

Bug number F ₁	F ₂	Metaphase I associations	Abnormalities	% fertility F ₂	F. parental fertility %
1 ₂	2	All cells normal	None		
	3	"	"		
	5	"	"		
	9	"	"		
	10	1 Fragment	1 Fragment		
2 ₁	6	All cells normal	None		
	7	"	"		
	9	"	"		
	11	"	"		
	15	"	"		
4 ₂	1	1 Fragment	1 Fragment	82.7	3.8
	2	1 Fragment	"	36.3	3.8
9 ₁	1	1 Fragment	"	20.7	2.6

Table 4-11. Chromosomal associations of metaphase I in testis squashes of 'F₂' male R. prolixus from F₁ male (app batch no. 25) × normal female. All chromosomes mentioned as chains except where stated.

Bug number		Metaphase I associations	Abnormalities	F ₁ fertility %	F ₁ parental fertility %
P ₁	P ₂				
6 ₁	1	6 II + 2 III + 2 I	2 Translocations		
9 ₅	1	All cells normal	None		
	2	8 II + 1 III + I	1 Translocation		
28 ₂	2	6 II + 2 III + 2 I		46	46
32 ₁	1	8 II + 1 IV			
37 ₃	4	8 II + 1 III + I			
	7	8 II + 1 III + I			
	9	8 II + 1 III + I			
	10	All cells normal			

Table 103. Chromosomal associations at metaphase I in testis squashes of male *S. prolixus* from
F. female (egg batch No. 2) X normal male. All multivalents associated as chains except
chain 22121.

Bug number		Metaphase I associations	Abnormalities	¹ F ₂ ¹ fertility %	F ₁ parental fertility %
F ₁	F ₂				
8 ₄	1	8 II + 1 IV	1 Translocation		
	3	8 II + 1 III + I + ?	1 Translocation + ?		
	2	8 II + 1 III + I	1 Translocation		
	3	8 II + 1 III + I	"		
2 ₁	6	8 II + 1 III + I	"		
	3	8 II + 1 III + I	"		
	6	8 II + 1 III + I	"		
	7	All cells normal	None		
6 ₂	1	6 II + 1 IV + 1 III + 1/8 II + 1 III + I	1/2 Translocations	11.3	4.5
	2	6 II + 1 IV + 1 III + 1/8 II + 1 III + I	1/2 "		
	4	8 II + 1 IV	1 "		
	5	All cells normal	None		
7 ₄	7	All cells normal	"		
	10	8 II + 1 III + I	1 Translocation		
	17	8 II + 1 III + I	1 "		
	1	All cells normal	None	96.6	1.6
14 ₃	10	All cells normal	"		
37 ₆	11	1 Fragment	1 Fragment		
	13	1 Fragment	"		
	17	1 Fragment	"		

..... Continued.

Table 4.17. (Continued)

Seg. pairing		Metaphase I associations	Abnormalities	'F ₂ ' fertility %	F ₁ parental fertility %
P ₁	P ₂				
P ₁	2	All cells normal	None	100.0	3.7
	5	8 II + 1 III + I	1 Translocation	4.0	8.3
	3	4 II + 3 III + 3 I/6II + 2 III + I	2/3 "		
P ₂	6	6 II + 1 IV + 1 III + I	2 "		
	7	8 II + 1 III + I + F	1 Translocation 1 F		
	1	8 II + 1 III + I	1 Translocation		
P ₃	2	1 Fragment	1 Fragment		
	7	8 II + 1 III + I	1 Translocation		
	8	6 II + 1 IV + 1 III + I	2 "		
	10	8 II + 1 III + I	1 "		
	13	1 Fragment	1 Fragment		
	14	8 II + 1 III + I	1 Translocation		

F = Fragment

Table 4.24. Chromosomal associations at metaphase I in testis squashes of 'F₂' male R. prolixus from (1) F₁ male (egg batch No. 5), X normal female and (2) F₁ female (egg batch No. 3) X normal male. All multivalents associated as chains except where stated.

Bug number		Metaphase I associations	Abnormalities	F ₂	
F ₁	F ₂			fertility %	parental fertility %
13	2	All cells normal	None		
	3	All cells normal	"		
	4	All cells normal	"		
	5	All cells normal	"		
20	1	1 Fragment	1 Fragment	8.2	8.2
	2	8 II + 1 III + I	1 Translocation	23.9	8.2
28	2	6 II + 2 III + I	2	2.8	3.5
	3	8 II + 1 IV	1 "	19.8	3.5
20	1	8 II + 1 III + I	1 "	11.7	4.9
	2	All cells normal	None	100.0	4.9
	3	1 Fragment	1 Fragment		4.9
18	1	8 II + 1 III + I / Fragments	1 Translocation + F	41.7	2.2

F = Fragment

Table 4.23. Chromosomal associations at metaphase I in testis squashes of 'F₂' male S. prolixus from F₁ male (egg batch No. 4) X normal female. All multivalents associated as chains except where stated.

Bug number F ₁ F ₂		Metaphase I associations	Abnormalities	'F ₂ ' fertility %	F ₁ parental fertility %
13 ₁	1	All cells normal	None	100.0	15.5
	2	8 II + 1 III + I	1 Translocation	94.2	15.5
40 ₃	2	8 II + 1 III + I	1 "	29.7	26.6
	3	All cells normal	None	97.8	26.6
	4	All cells normal	"	100.0	26.6
	5	All cells normal	"	"	"

Table 4.26. Summary of results from Tables 4.20 - 4.25 to show chromosomal abnormalities of 89 'F₂' male R. prolixus from 4 parental egg batches.

Abnormality	Parental egg batch								Total	%
	1		2		3		4			
	♂	♀	♂	♀	♂	♀	♂	♀		
None	8	9	2	6	5		4		34	38.2
1 Fragment	1	4		5	2				12	13.5
1 Translocation	7		5	16	3	1	2		34	38.2
2 Translocations	1		2	4	1				8	9.0
3 Translocations				1					1	1.1
Total	17	13	9	32	11	1	6	0	89	100.0

♂ = Progeny of F₁ ♂ X normal ♀

♀ = Progeny of F₁ ♀ X normal ♂

individual bugs, although the most frequently encountered orientation assumed by the chain was linear, i.e. parallel to the long axis of the spiracle. As with P_1 males, some P_2 spermatocytes contained in the same cell the multivalent could assume different modes of orientation as in Plate 4.12 which shows a spermatocyte from P_2 male No. 5₁₁ with 2 translocations associated as 6 II + 1 IV chain & a III chain (7), the chain of IV showing parallel orientation and the chain of III linear orientation. In examining the squashes from testes of all these ' P_2 ' males not a single ring association multivalent was seen. These results suggest that the majority of heritable translocations passed on from P_1 parents to ' P_2 ' males were probably small interchanges producing sygotene configurations with one small arm, which would be undetectable, producing the bulk of III + I associations in these bugs.

The results show that in 12 of these ' P_2 ' males, the only detectable chromosomal abnormality was the presence of one or more chromosomal fragments. Plate 4.11 shows a metaphase I spermatocyte from ' P_2 ' male No. 25₁₁ with a single fragment and Plate 4.12 shows a chromosome spread from P_2 male No. 5₁₁ with 2 chromosomal fragments. The presence of these fragments in meiotic cells of ' P_2 ' males is an important finding in relation to the structure of *H. pallidus* chromosomes. If *H. pallidus* were monosomic, with half the normal chromosome complement, they must since they would have been acentric and therefore unable to attach to the spindle; the persistence of these fragments to the ' P_2 ' generation offers good evidence for the holokinetically nature of triatomine chromosomes.

The results of the cytogenetic examination of the ' P_3 ' bugs raised in this experiment are shown in Table 4.27. These results show that none of the 11 ' P_3 ' males examined showed any detectable chromosomal abnormalities.

Table 4.27. Results of cytogenetic examinations of 11 'P₃' male R. prolixus descended from P₁ males of the first parental egg batch.

Egg type		Chromosomal abnormalities	Percent of Metaphase II	Percent of Metaphase II
P ₁	P ₂			
2 ₁₊₁	1	normal	100.0	100.0
	2	—	—	—
2 ₂₊₁	1	—	100.0	—
	2	—	—	—
	3	—	—	—
	4	—	—	—
	5	—	—	—
3 ₁₊₁	2	—	6.0	3.1
2 ₃₊₁	1	—	41.9	3.8
	2	—	—	—
	3	—	—	—

Correlation of fertility and chromosome anomalies

The F_1 generation of M. trilineatus bred from irradiated male parents and normal female parents had a much lower mean fertility than their parents when they were crossed with normal males or females or approximately, and cytogenetic examination of the testes of the F_1 males showed that each was carrying at least one translocation. The 27 F_1 males with a single translocation mated with normal females resulted in 1718 eggs, 150 of which hatched successfully giving a mean fertility of 8.7%. The 18 F_1 males shown to have 2 translocations when mated with normal females resulted in 1151 eggs, 69 of which hatched giving a mean fertility of 6.0% and the 4 F_1 males with 3 demonstrable translocations when mated with normal females resulted in a total of 113 eggs none of which hatched. Although there were no demonstrably significant differences between the mean values for fertility of males carrying 1 or 2 translocations, it is more informative to examine the proportions of large in these 2 groups which were completely sterile. From the data presented in Tables 4, 5 & 4, 18 it can be seen that of the 27 F_1 males with a single translocation, 12 (44.4%) were completely sterile and of the 18 males with 2 translocations 17 (95.6%) were sterile, and that all of the large with 3 translocations were completely sterile. These results do suggest a correlation between degree of chromosomal abnormality and fertility.

The fertility results for F_1 males show that their fertility may be related to their histone content. Those males from which potential egg batches being significantly more fertile than those from which no egg batches. The cytogenetic results presented in Table 4, 18 show that the number of detectable chromosomal abnormalities increased with F_1 male histone content. Thus, from the first potential egg batch it resulted males were normal, and 6 had one translocation each and 2 had 2 translocations. Examination

of the other egg batches shows that males from later egg batches had greater degree of chromosomal abnormality and this is closely related in their reduced fertility and in the increased proportions of completely sterile matings.

F_1 male bugs were found to be significantly less fertile than their females. Males crossed with normal females produced 12.64% fertile progeny, while females crossed with normal males produced 44.6% fertile progeny. Since only males could be assigned for chromosomal abnormalities, a cytogenetic origin cannot be readily ascribed to these differences between male and female F_1 bugs in fertility.

The 28 ' F_1 ' bugs crossed with normal mates had a mean fertility of 44.6% which was much greater than the mean fertility of the F_1 generation of 12.64, although only 28 selected F_1 bugs were tested, 27 males and 1 female cytogenetically. These proportions of fertile progeny from chromosomal abnormalities in the bugs of this generation were much reduced when compared with the abnormalities of the F_1 males, for while only F_1 males crossed with normal mates, only 28.6% of F_1 males had detectable chromosomal abnormalities. Moreover, 3 of the 28 F_1 males which were crossed with normal females were fully fertile and examination of their testes revealed no detectable abnormalities. These 6 bugs were numbered 14_{3,1}; 37_{7,2}; 28_{5,2}; 13_{1,1}; 40_{3,1} and 41_{3,1}, the first 2 being females produced from F_1 ♀ X normal ♂ crosses, and, therefore, completely cytogenetic. Data on fertility are given in the Table. Moreover, 15 other 4 males were produced from F_1 ♂ X normal ♀ crosses and the results show that the father of 28_{5,2} had detectable translocations and the fathers of the other 3 males had a single translocation. These results suggest that the translocations of these 4 F_1 parental males were inviable and were not transmitted to their progeny.

Only 2 of the 28 ' F_1 ' bugs crossed with normal mates showed evidence of having reduced fertility in comparison with their parents and these

were bugs Nov. 30, 1961, which showed evidence of carrying a single translocation, and 28, 1962, which had 2 translocations, their parents showing configurations which indicated 1 and 2 translocations respectively. The apparent stability of these translocations over 2 generations suggests that the chiasma association found in these bugs were capable of convergent orientation to produce viable gametes. Unfortunately, these families were not crossed to produce further generations, so that the further stability of these translocations was not tested.

It has been shown that the mode of orientation of chain multiples is critical in determining their viability and, although these orientation patterns were not quantified, it appears from the photomicrographic evidence that the majority of the P_1 males showed multiples which were orientated linearly which would inevitably lead to low fertility, and many of the translocations would not be passed on to the next generation. The cytogenetic results are in accordance with this, in that while all P_1 males showed some translocated chromosomes, 80% of the F_2 males showed no evidence of any translocation. Furthermore, only 2 of the 28 F_2 males showed 1 or 2 copies of 1 translocation, compared with the 28% of P_1 males with 1 translocation. This result probably reflects the increasing improbability that a multivalent would align in a viable mode in the same cell.

The cytogenetic results also demonstrate that even a single chromosomal fragment can have a wide ranging effect on the fertility of the male carrying it; thus P_1 male $2n_{24}$ had 4 chromosomes in pairs and a fertility of 84.7% while P_1 male $2n_{25}$ had 1 chromosome with a single fragment, was infertile, completely sterile, and F_2 male $2n_{26}$ with 2 fragments (one with 1 and 2 fragments) had a fertility of 81.7%. In the

case of $P_2 \times 28_{S_{+3}}$, the single fragment clearly had the effect of a dominant lethal mutation.

The overall increase in P_2 fertility has been seen to be a reflection of a decrease in the amount of detectable chromosomal abnormality in the males of that generation. Namely, cytogenetic examination of the P_2 males provided no explanation for the superiority of the P_2 female over the P_1 male in terms of fertility. The results in Table 4.26 show that, in the first egg batch, 8 P_1 males from $P_1 \times X$ normal crosses carried translocations while none of the males from the $P_2 \times X$ normal crosses showed any translocations. These findings were reversed in the second egg batch in which males from $P_1 \times X$ normal crosses showed the highest translocation frequencies.

Only 14 P_2 males, all descended from the first parental egg batch, were produced for cytogenetic examination and none of these showed any signs of detectable chromosomal abnormality in their spermatocytes. It may reasonably be assumed that their fertility would have been nearly normal had it been assessed. P_3 male No. $3_{P_{+1}}$ is of special interest in this group in that its male parent P_2 $3_{P_{+1}}$ had a single translocation and fertility of 8.3%, and its male grandparent P_1 4_2 had 7 translocations associated with a fertility of 1.3%. This is a clear demonstration of how a bug with a large amount of chromosomal abnormality can, by 1 generation of out-crossing to the wildtype, produce progeny without any detectable abnormalities in their meiotic chromosomes.

DISCUSSION

The dose-response curve produced as a preliminary requirement for this experiment revealed differences, both in amount of radiation required for complete sterility and in rate of response, between the curves

produced for the production of complete sterility in adult male *D. melanogaster* by Shaver (1963) and in the present work. Much higher doses were required by these workers to produce complete sterility in adult male *D. melanogaster* and the reason for this most probably lies in the different radiation source used. Kligerman and Searle (1968) have shown that there is a qualitative difference between γ -rays and X-rays of the same dosage; they found X-rays to be twice as effective as 60°Co γ -rays of the same energy in producing chromosomal aberrations in *D. melanogaster*. Similarly, Searle et al. (1968), in studying translocation frequency in mouse spermatogonia, found that at low dose rates X-rays induced twice the frequency of translocations as did γ -rays of the same energy while at high doses this difference was reduced. Although interpretation of these results is difficult, Searle et al. (1968) concludes that the kinetics of translocation induction was different for the 2 types of irradiation. The differences in dose-response curves demonstrated here serve to confirm that variation in dose-rate and radiation source can produce widely differing changes in the same organism.

D. melanogaster males irradiated with 6 K rad γ -rays were crossed with untreated females. Offspring of these F_1 progeny were screened for normal males. The resultant F_2 males were then screened for normal males. Fertility and fecundity were measured for all specimens. Representative F_1 , F_2 and F_3 males were examined cytogenetically and the results of these studies of meiotic preparations were compared with the fertility data. The results show that the reduction in fertility of the 2 generations of males correlated well with the degree of abnormality demonstrable in their spermatocytes. The most frequent chromosomal abnormalities encountered were translocations and, in the F_1 generation, males carrying the greatest number of translocations were generally less

fertile. The maximal number of translocations seen in F_1 males was 2 and all bugs carrying this number were completely sterile. These results are similar to those obtained by Bauer (1967) and LeChance et al. (1970) working with other holokinetid insect species. The high recovery rate of translocations in the F_1 males in this experiment is in agreement with Bauer's (1967) theory that dicentric chromosomes cannot be produced in holokinetid species, so that all forms of interchange should be viable.

When mated with normal bugs, the F_1 generation, *h. transloc.*, whether male or female, resulted in lower fertility of eggs than their nontranslocated parents crossed with normal females. This finding suggests the 'haploid sterility' demonstrated by several workers in eurytomids. For example, Korte and Hildebrandt (1968) showed that F_1 males although fertile were more sterile than their fathers which had been given sub-sterilizing doses of radiation. However, LeChance et al. (1970) found no delayed sterility effect in their experiments with irradiated diamond-back.

The cytogenetic evidence from the present work has shown that the greatly reduced fertility of F_1 males may be related to the form of association and resultant orientation difficulties of multivalents in these translocation heterozygotes. The majority of interchange associations, at first metaphase, as a chain trivalent (III) plus a univalent (I), which must have arisen due to the failure of chiasma formation in 2 adjacent arms of the syntelic pairing bivalent during meiotic prophase. Chains of IV were also commonly found in F_1 interchange heterozygotes. This presumably arose because of the failure of a single chiasma. Lyon and Marston (1965) in a study of translocations in mice, found that translocations producing sterility showed a higher frequency of chain associations than did those which did not lead to sterility. The high frequency of multivalent chain associations seen in F_1 males in the

present experiment was associated with a very high degree of sterility (84.54 per cent sterility) calculated from the proportions of eggs which hatched. The genetic consequences of chain formation depend on the mode of orientation of the multiple in relation to the spindle axis, in a metacentric species, it would be expected that the linear, convergent, indifferent and parallel modes would be assumed with equal frequency. However, M. troglorum normally shows linear orientation of the multiple bivalents (Bock, 1967) with axes aligned to the long axis of the spindle, so that it would be expected that linear orientation would be favoured by chain multivalents. The results show that this is the case; most of the chain multiples seen in the present study were linearly orientated. Lewis and John (1963) have shown that only the convergent orientation of chain multiples allows the production of balanced gametes, other modes leading to duplication and deficiency of genetic material and culminating in embryo death. The small reduced fertility of M. troglorum found in this experiment is therefore in agreement with both the cytological evidence and theoretical expectations.

Lawrence et al. (1971) demonstrated the preferential production of chain multiples in the bug A. pallidus, but did not comment on the orientation of these chains; judging from their published photographs, the multiples showed a mainly linear orientation to the translocation heterozygotes. The present work has shown that, although linear orientation of chain multiples is favoured in M. troglorum, the 3 other possible modes of orientation (convergent, indifferent and parallel) were also assumed, if not in frequency, indeed, in individuals heterozygous for more than one translocation, a mixture of orientation modes was observed. Since at least one such case (1971) and only one interchange heterozygote was found in this experiment with a

demonstrable ring associations in meiosis, but this result is important because it shows that although such associations are exceptional, their formation is not precluded in tetrahymena eggs. Bauer (1967) found that ring multiple associations were most common in lepidopteran interchange heterozygotes, only rarely being multiple being found. This striking difference between organisms with apparently similar holokinetic chromosomes must be partly a function of chromosome morphology. The apparent infrequency of ring formation in H. p. mixta suggests that most chromosome breaks were small, producing fragments which almost never met, since chiasmata frequency in H. p. mixta is normally low, only one chiasma per bivalent being formed (Uehelaw, 1966), the formation of chains is likely to be encouraged in this bug. Jais and Jones (1965) suggest that the genotypic properties may also influence the type of orientation of interchange multiples formed in different organisms which may account for the differences between the present results and those of Bauer (1967) and LeChance et al. (1970).

The fertility studies of F₁ H. p. mixta showed a significant linear trend downwards in fertility, related to the birth order of 4 separate batches of F₁ bugs, which had been reared from 4 successive egg batches laid by their mothers after mating to randomised partners. Cytological examination of the F₁ males from these 4 successive egg batches revealed that the males from the later egg batches with lower fertilities also showed the greatest amount of chromosomal abnormality. If it is accepted that these insects do not suffer the effects of parity that a viviparous animal would, then these chromosomal differences would most likely account for the major part of this reduced fertility. The increased chromosomal damage of males of later egg batches may be explained by reference to the experimental design by which the parental females were

fed 4 times at 14 day intervals, the eggs being collected to form one egg batch prior to the next feed. It is known that adult M. proluxus of both sexes mate repeatedly (Baldwin and Shaver, 1963) and therefore the first egg batches were probably fertilized by sperm which were mature spermatozoa at the time of irradiation, while the later egg batches were probably fertilized by sperm which at the time of irradiation may have been spermatocytes or spermatogonia. This being so, the chromosomes of prophase spermatocytes or spermatogonia would be in a less condensed state than those in the sperm head and would have been subject to more hits, which would increase the frequency of breaks and interchanges. This could account for the differences in fertility of the 4 batches of males reared from eggs laid over a period of 8 weeks after the initial irradiated parent male X normal female crosses.

An interesting feature of the analyses of fertility data for M. proluxus was the superior fertility of F_1 females when mated to normal males, over that of F_1 males mated to normal females. Unfortunately, cytological examination of ' F_2 ' males raised from these 2 types of crosses revealed no obvious cytogenetic cause for this result, and cytogenetic examination of F_1 females was not possible. Snell (1946) in a study of radiation-induced translocations in the mouse found that for 5 of the 6 induced translocations, males had a higher fertility than females carrying the same abnormality. Join and Lewis (1965) sought to show that this difference was due to the reduced frequency of multivalent disjunction in females because of their higher chiasma frequency. However, it had been assumed that chiasma frequency was greater in female mice because of the higher recombination frequency in that sex (Green, 1966), and it has since been shown (Henderson and Edwards, 1968) that chiasma frequency is in fact higher in female mice than in males. It

would appear that chiasma frequency may account for this result in mice, but as the chiasma frequency of female R. prolixus is not known, it is not possible to relate this character to the present results. John and Lewis (1965) have further suggested that the time available for orientation of a multiple could influence the mode of orientation taken up, and that this was perhaps a genotypically related factor. It may be that the time available for multiple orientation is greater in the egg than in sperm of R. prolixus because of the much slower rate of egg production compared with sperm production, and that this allows time for more genetically stable egg nuclei to be produced in females. However, in the absence of further cytological information, this explanation can only be speculative.

The increased fertility of ' F_2 ' males compared with that of their fathers was found to correlate with cytogenetic differences between the 2 generations. Thirty-four of the 89 ' F_2 ' males examined showed no evidence of any chromosomal abnormality in their spermatocytes, and since every F_1 male examined had at least one translocation, this reduction in observable chromosomal aberration must account for the superiority of the ' F_2 ' bug fertility. This would suggest further that it is difficult for an interchange to persist from one generation to the next in this species, probably because of the holokinetic nature of its chromosomes linked with a predilection for axial orientation and a low chiasma frequency. However, 43 ' F_2 ' males did show evidence of carrying translocations suggesting that linear orientation of chain multiples is by no means compulsory in this system and that it may be possible to establish a heritable interchange in a population of bugs.

Several F_1 males, as well as exhibiting translocations, showed chromosomal fragments in their spermatocytes and 12 of the ' F_2 ' males examined showed no other chromosomal abnormality than a fragment. The

persistence of such fragments through 2 generations is evidence of their mitotic and meiotic stability and provides proof of the haplokinetic nature of K. prolixus chromosomes which has up till now been inferred from ultrastructural studies (Black, 1967). Transposons were found to have a variable effect on the fertility of males carrying them, so that a male with a single fragment may have been completely sterile while others with 2 fragments resulted in approximately 50% sterility. Lawrence et al. (1968) found that 3 fragments would induce complete sterility in the milkweed bug, but this was not found to be true in K. prolixus. It seems that the relative importance of fragments may depend more on the degree of genetic importance of that part of the genome incorporated within them, their action being that of dominant lethal mutations by either genetic duplication or deficiency.

None of the 11 P_1 males examined showed any evidence of carrying any cytogenetic abnormalities, suggesting that the transposons present in their parents' spermatocytes were not stable. However, it may be presumptuous on the basis of this limited number of observations to assume that this would be the fate of all translocations induced in tritumid bugs. For example, P_{21} males (1967) with chromosomal interchanges had only slightly reduced fertility compared with their parents (these lines P_{11} and P_{12}) and it may be that some particular translocations were relatively stable; but unfortunately these bugs were not bred by a further generation so this must remain a speculation.

Since K. prolixus is an important vector of Chagas' disease in Venezuela, the results of this experiment must be considered in addition to the control of this pest by genetic means. Cummings et al. (1965, 1964) and Baldwin and Chant (1970) ruled out the use of sterile male release as a control measure for K. prolixus populations because of the

high radiation doses required to completely sterilize males, which resulted in bugs of greatly reduced fitness. Haidich and Chant (loc. cit.) experimented with irradiation of *M. polius* in an atmosphere of nitrogen and found that this improved the fitness of those bugs compared with those irradiated in air. Because of the high radiation doses of the chromosomes of *M. polius* demonstrated in the present work, it would seem appropriate to assay control measures which take advantage of the peculiarities this confers on trisomic bugs, advantages which have been usefully employed in controlling Lepidoptera, since the 'delayed sterility' demonstrated here for *M. polius* given sub-sterilizing doses of Y-rays, is an effect characteristic of holokineton breaks and first observed by Provata (1962) in the treading moth, *Plutella maculipennis*. A further useful feature of giving sub-sterilizing doses of radiation to Lepidoptera was found to be a distortion of the sex-ratio in the F_1 so that the majority were males (Haidich, loc. cit.). This sex-ratio distortion in favour of males which is so desirable for genetic control programmes has also been demonstrated in irradiated *Trichoplusia ni* by Partin et al. (1971) and suggested that if sex is related to the extra large X-chromosome of female-determining sperms being more susceptible to radiation damage. The present work has shown that irradiation does not distort the sex-ratio of the progeny in *M. polius* and this may be due to the very small size of the sex chromosomes in this species.

Despite this drawback related to chromosome size, the present work has shown that sub-sterilizing doses of radiation will induce delayed sterility in *M. polius* and, as Knippling (1971) has shown for Lepidoptera, it would be more effective in controlling populations of this species if partially rather than fully sterile males were employed. Although

K. trilineatus may mate repeatedly, this may not be a disadvantage as von Borstel (1966) has pointed out that monogamous matings are not essential for the success of sterile male control programmes, for the true competition is between normal and lethal sperm within the female spermathecae and it is therefore irrelevant how many times an insect mates.

The present experiment has demonstrated that while the linear orientation of multivalents is favoured in K. trilineatus, leading inevitably to the loss of translocations, it is possible for certain multivalents to orientate in a stable fashion and also for ring multivalents to be formed in this way. A long-term association between could perhaps produce a translocation which preferentially orientated in a stable way during meiosis, but the long association of translocation bursae, prior to the an obstacle to such a scheme.

GENERAL DISCUSSION

The triatomine bugs of South and Central America are of great medical importance as vectors of Chagas' disease which seriously affects the lives of some millions of people. It is therefore appropriate that the results of the present work should be considered in relation to the problems associated with reducing the incidence of this disease. The elimination, or reduction in size of, vector population whether triatomines or vectors of pathogens other than 2. *Trypanosoma cruzi* is of obvious use in reducing the incidence of a disease.

Insecticides, particularly indoor residual deposits of contact insecticides such as DDT and other chemicals more recently developed, revolutionized, and have progressively advanced, methods of vector control for man, insect-borne diseases over the past 30 years. For triatomines, indoor residual deposits of organochlorines (BHC and dieldrin) in rural and urban housing throughout much of the South and Central America have been particularly successful in eliminating or reducing domiciliary infestations of vectors of Chagas' disease and where financial resources were adequate, improved housing standards have also contributed to control of the vector and the disease. Re-infestations of treated premises do, however, occur with species which have, in part, a 3. *Triatominae* habit of being indoors, or due to accidental re-introduction of bugs.

In general, medical usage of toxic insecticides is 4. *insecticides* much less contaminative of the general environment than methods used in agriculture such as aerial spraying of crops, although mal problems do sometimes necessitate outdoor sprayings of this kind. 5. *DDT* Reinfestations of treated premises by 6. *Trypanosoma cruzi* 7. *Triatominae* control methods, the polluting effects of insecticides have attracted considerable criticism in recent years. As a result, attention has been directed to genetic

methods for pest and vector control, which could circumvent hazards to human and animal health from insect resistance. It may also be said that by insecticide control elimination of a vector is seldom complete, although often providing reduction in population density consistent with virtual eradication or effective reduction in the incidence of transmission of disease and pest of livestock. Results of this order have been achieved over large areas against *Simulium*, notably *S. vittatum* and *S. morsitans*, against *Culex* mosquitoes, notably *C. quinquefasciatus* and *C. tritaeniorhynchus*, and against *Anopheles* mosquitoes, notably *A. gambiae* and *A. stephensi*. Reduction of these species has yet been controlled thoroughly and, moreover, indications of insecticide resistance in *S. vittatum* and *C. tritaeniorhynchus* provide a warning that, as in other insect pests and vectors, this serious consequence of repeated chemical treatment may become a practical problem. Other control measures warrant investigation on this account alone. Genetic methods of control necessitate thorough investigation at laboratory level if they are to be adequately evaluated on the basis of sound scientific knowledge. However, before the principles of genetics can be applied to the regulation of insect populations, as much information as possible must be gathered about the whole biology of the pest or vector species in question. *Anopheles* and all species of mosquitoes are a good example of vectors. For example, the commoner mosquito *Anopheles gambiae* has complex genetic make-up, consisting of two distinct sub-species, although the adult forms are indistinguishable, and while some strains are known to act as malarial vectors, others were not. By crossing these strains, Hackett (1937) found that hybrid crosses were sterile thus demonstrating that distinctive sub-divisions of this species existed. This resulted in an *Anopheles gambiae* complex. It

is clear from this example that taxonomy, even at the egg-stage, can play an important role in studies of vectors. More recently, an elaboration from the classical criteria of morphology, has been the highly satisfactory technique of chromosome banding. In the case of the Trichotomus complex, Trichotomus trichotomus and Trichotomus trichotomus have been identified by the analysis of the species complex, other stages remained morphologically indistinguishable until it was shown that the salivary gland chromosomes of the larvae provided an excellent means of differentiating the members of the complex (Columbi and Sabatini, 1967). Trichotomus and Trichotomus larvae salivary glands showed complex banding patterns when stained with Giemsa, and the bands were found to be highly stable. Cytogenetic taxonomy was also developed to differentiate members of the Trichotomus complex, in which the larvae were identified as vectors; the discovery of polytene chromosomes in the nurse cells of the ovaries of the adult females of these mosquitoes being of particular convenience for identification of adult members of the 6 sub-species (Columbi and Sabatini, 1967).

Hybrid sterility studies of the Trichotomus have been made, and indeed are still being carried out (Perlovskaya-Szumlewick, 1974), but such genetic experiments are very slow because of the lengthy life-cycle of the bugs. Furthermore, unlike mosquitoes, the immature stages possess polytene chromosomes to enable in situ banding. The present study of polytene chromosomes in Trichotomus bugs confirmed that feeding animals, chromosomes are not becoming visible as in the case of feeding, the only differentiating feature between certain species being a banding pattern in the 1st and 2nd chromosomes. The cytogenetics of the Trichotomus appeared to be an intractable problem

language of these presumed chromosomes. The chromosomes have been improved during the researches of this thesis by application of techniques of chromosome differentiation developed for mammalian chromosomes. It is now the chromosomes of Trichogramma which are large, which are stained well, when stained by the Feulgen method. The chromosomes are stained by the use of Feulgen for differentiating species. The patterns of banding for Trichogramma and Nicrophorus which are species specific. This banding technique deserves to be investigated for a wider range of Trichogramma species. The chromosomes are similar to tritome taxonomy some of the precision while the study of polytene chromosomes has given to the study of mosquito species and crop as of medical importance.

The present study of spermatogenesis in Trichogramma is the second part of this thesis. The chromosomes and chromosomes and chromosomes affect the rate at which spermatocytes develop. In particular, the process of spermatogenesis has been shown to be inhibited by starvation in fifth instar larval males, and it was found that spermatocytes spend longer periods in the 'diffuse stage' of meiosis during such periods of starvation. Although this particular aspect of the study was not directly related to the chromosomes of Trichogramma, it is nevertheless worthwhile to consider these results in relation to this problem. The results show that meiosis and spermatogenesis are chromosomes by the chromosomes of the chromosomes. The chromosomes of the chromosomes are not clear, except that it is linked in some way to the nutritional status of the bug. If further research revealed that this control mechanism was chemical, then it is possible that the process of

spermatogenesis could be inhibited to provide an alternative method of sterilizing either wild populations, or laboratory-reared or wild-caught males for a sterile-male control programme.

The third part of this thesis examined the genetics of bug susceptibility to infection in vitro, a study which has obvious applications since xenodiagnosis still occupies a central role in the clinical diagnosis of chronic Chagas' disease. A highly susceptible strain of tritarsine bug would improve the sensitivity of this diagnostic test, and also fewer bugs than the 10 bugs used on each of the 4 occasions customarily recommended at present for each patient tested would be required to be fed to each patient, thereby decreasing the discomfort and, in some individuals, severe sensitization which this test feeding inevitably causes for the patient. The results of the present study have shown that the intensity of faecal infection of individual bugs is, in part, genetically determined. Moreover, on the basis of quantitative differences in faecal infection, produced in these generations between discrete populations of R. prolixus, one group excreting significantly greater numbers of trypanosomes than the other, indicating differences in susceptibility to infection. The results also revealed a most interesting difference between sexes of R. prolixus, males excreting significantly greater numbers of trypanosomes than females, despite the fact that females ingested more blood (and thereby trypanosomes) of the infected host, thus leading clearly to the suggestion that faecal infection in bugs is determined by the amount of trypanosomes ingested. For example, Bittencourt et al. (1973) related the percentage of bugs with positive faeces at xenodiagnosis to the amount of blood ingested by the bug, but the present work confirms the conclusion of Bertram and Phillips (1967) that the numbers

Furthermore, Penetration Experiment Group II 1961 have recently shown that *Paratransferrin* super-numerary larvae, produced by juvenile hormone treatment, had much lower infection rates than control bugs when fed on chronic Chagas' infected patients, despite the fact that they did not exhibit any significant difference in the infecting fever; this supports the present conclusion, that blood-meal size is not directly related to subsequent fascial infection in bugs. The present experiment demonstrated that it is possible to breed bug with increased susceptibility to *T. cruzi* infection, and it is suggested that inbred populations of susceptible *R. prolixus* should be bred for use in xenodiagnosis and preferably only males of such colonies should be used.

The results obtained by irradiating male *R. prolixus*, mating them with normal females and following the results through successive generations of out-crossing, must be considered in relation to the control of wild populations of this vector species. The results have demonstrated that sub-sterilizing doses of γ -irradiation produce F_1 generation offspring males with reduced fertility compared with their irradiated fathers. This delayed effect of sterility has been related, by parallel studies of chromosomal abnormalities, to the holokinetic structure of bug chromosomes, and is similar to the situation encountered in lepidopteran pests. Knippling (1970) has suggested that where an insect species demonstrates a high level of radio-resistance, control is more effective when semi-sterile rather than completely sterile males are released. Two reasons are cited: Firstly to sterilize fully the males would make them unfit and unable to compete with normal males at mating and, secondly, the delayed effect on

fertility of releasing semi-sterile males would produce a longer-term suppression of population numbers. In the present experiment, however, the fertility of most of the progeny of semi-sterilized male P. prolixus was returning to normal by the second generation of outcrossing to normal males. Thus, the sterility substerilizing does on wild populations of P. prolixus may be relatively innocuous. However, the practical dynamics which would arise in such a situation must be taken into account in terms of repeated releases into a population of bugs already partially translocated or translocated. Certainly, a sterile male release program would be the basis of longer duration and its vectors have social consequences and constraints. Since male P. prolixus are, like females, translocated vectors of Yagui's disease, the release of large numbers of males in a control project would not be socially acceptable to the people living in endemic areas, nor, with a disease of such severity, highly dependent by medical intervention. It is more feasible to suppose that such releases could be justified to replace the natural population in an area with one carrying a translocation linked to, say, genes for insecticide susceptibility in the event of insecticide resistance becoming a serious public health problem. This, although so far a matter of little practical consequence, may yet arise as a matter of social concern to the public health authorities. Conceivably, male releases, whether of semi-sterile irradiated males or males carrying translocations linked with insecticide susceptibility, should be more acceptable to medical authorities if these control attributes were also genetically coupled with use only of males of a strain minimally susceptible to infection with the pathogenic organism, Trypanosoma cruzi.

Clearly much still remains to be done in experimental genetics

tentative prospects can be elaborated to realistic appraisals of the potentialities of genetic manipulation in triatomines for the reduction or eradication of Chagas' disease by genetic control of its vector insects.

SUMMARY

PART ONE

1.1. A study of meiosis in males of 12 species of triatomine bug (Triatoma brasiliensis, T. infestans, T. lecti, T. maculata, T. tibio-maculata, T. phyllasma, T. protracta, T. vittigena, T. pseudomaculata, T. sordida, Rhodnius neglectus and R. prolixus) from South and Central America showed that the basic complement for this genus was 20 autosomes + XY. The greatest amount of variation between species was in the number of X-chromosomes which varied from X₁Y to X₁X₂X₃Y. These X_nY systems are thought to have originated by fragmentation of an archetypal X-chromosome. One further species, Panstrongylus megistus was however found to only have 18 autosomes + XY.

1.2. Female meiosis was observed in a single species, R. prolixus, the first maturation division in an egg being observed about the time of ~~oviposition~~ oviposition.

1.3. A technique is described for the production of G-bands in the chromosomes of ~~monokaryotic~~ from triatomine embryos. The results obtained with R. prolixus and T. infestans suggest that the technique will provide genetic markers of value in elucidating problems in the systematics, biology and genetic control of these two important vectors of Chagas' disease, and of other triatomine species of South and Central America.

1.4. Meiosis in F₁ males of the cross T. pseudo-maculata X T. sordida and in males from the reciprocal cross was found to be characterized by multiple associations and univalent formation. The orientation of the chain multiples made the survival of hybrid interchanges seem improbable, and the sterility of these F₁ bugs most probably related to these chromosomal events.

SUMMARYPART TWO

2.1. Examination of squash preparations from testes of fourth and fifth instar larval R. prolixus showed that testis differentiation would only occur in fifth instar larvae and it is suggested that some genetic switch mechanism associated with age may be necessary to stimulate testis differentiation and that juvenile hormone may be involved in this process.

2.2. It was found that spermiogenesis was inhibited in diapausing fifth instar larvae, the primary spermatids produced by these bugs becoming 'aborted' and probably removed by autolysis.

2.3. Following a histological, ~~pelvic dissection~~ and spermiogenesis were rapidly initiated in fifth instar larvae, mature spermatozoa being found in their testes prior to the moult to the adult.

2.4. The rate of meiosis was compared in fed and diapausing fifth instar larvae and adults by ^3H -thymidine labelling of spermatocytes. Meiosis occupied 21 days in unfed fifth instar larvae and 12 days in fed larvae. Similar differences were demonstrated among fed and unfed adult testes, the difference in time taken being largely accounted for by variation in the length of time spent by spermatocytes in the 'diffuse stage' of meiosis.

2.5. Introduction of ^3H -uridine into testes in ^3H -uridine demonstrated that RNA was being actively synthesized in 'diffuse stage' nuclei of both fed and diapausing testes. It is proposed that during the 'diffuse stage' the chromosomes, which are extended and may, possibly, have a functional significance similar to that of lampbrush chromosomes, could be engaged in the synthesis of messenger RNA in response to the nutritional status of the bug and its age, so governing the further stages of spermatogenesis.

SUMMARY

PART THREE

3.1. Susceptibility rate and intensity of infection were measured for 2 populations of R. prolixus selected over successive generations for susceptibility and refractoriness to 2 different strains of T. cruzi.

3.2. Selection for 2 generations with the Peru strain of T. cruzi, and for 3 generations with Strain 7 T. cruzi, failed to demonstrate continuous variation which could be ascribed to major gene differences controlling susceptibility rate in R. prolixus populations.

3.3. Selection for differences in intensity of infection in individual bugs produced the following quantitative changes in the experimental bug populations:-

i. Highly significant differences were found between similar populations of R. prolixus infected with 2 different strains of T. cruzi, suggesting that trypanosome genotype can affect the level of intensity of infection in the bug.

ii. Significant differences in mean values of intensity of infection were found among selected families of bugs infected with the Peru strain or Strain 7 T. cruzi, and for all generations raised. These family differences reflected a degree of resemblance between relatives characteristic of a continuously variable character under polygenic control.

iii. Analyses of male and female results revealed significant differences between sexes, males having consistently higher levels of intensity of infection than females, suggesting either sex-linkage or sex-limitation of the character.

3.4. i. Two-way selection for high and low levels of intensity of infection among bugs of the T. cruzi Strain 7 group produced, in the 7th generation, 2 clearly separated groups of families with significant differences in mean scores between families of the 2 selection lines.

ii. Response to selection when plotted against selection differential for selected families of the Strain 7 group of bugs produced highly significant regressions which were used to calculate the realized heritability for the characters. The heritability for intensity of infection measured from the divergence of the two regressions of response on selection differential was found to be 5.5%.

iii. The response to selection was asymmetrical being twice as great in the downward direction (increasing refractoriness) as in the upward direction (increasing susceptibility). It is expected that this asymmetry may have been caused by the dominance of genes for increasing intensity of infection and/or by inbreeding depression of a fitness-related character.

1.5 It is expected that the efficiency of selection may be improved by the production of an inbred line of *K. prolixus* selected for high intensity of infection with *T. cruzi* and that only male bugs of such a population should be used for selection.

DISCUSSION

PART FOUR

4.1 Eggs from crosses of 40 adult male *K. prolixus* irradiated with 6 K rad γ -rays with normal females had a mean fertility of 23.9%, only 2 being completely sterile. The mean fertility for control eggs of normal males was 11.8%. The 50 F_1 progeny of the irradiated male 2 crossed female parents had a mean egg fertility of 17.8%, and 43 of these crosses were completely sterile.

4.2 Twenty-eight F_2 bugs raised from F_1 2 normal females were mated with normal partners and had a mean fertility of 44.6%, 6 of these having fully fertile, a normal female cross fertility.

4.3 These findings of initial reduction of fertility followed by recovery of fertility by the 'P₂' generation, following irradiation of the males of the original parental generation (*domes*, *chromosomal*), changes in chromosomal configurations as observed by cytogenetic examination of preparations of male testis.

4.4 The survival to the 'P₂' generation of chromosomal fragments confirmed the holokinetic nature of tritamine chromosomes.

The very high recovery rate of translocations in P₂ generation males can be related to the holokinetic chromosome system of these bugs which precludes the formation of dicentric chromosomes which are inviable in monocentric species.

4.5 In P₂ and P₃ males the majority of translocations were associated in chains of III + 2 or in chains of IV. Only one bug was found with a ring of IV chromosome association and it is suggested that chromosome morphology combined with a low chiasma frequency hinders chain association formation.

4.6 Most chain multivalents showed linear orientation which may lead to segregation deficiencies and apogametes. Survival of translocations in *H. prolixus* will be difficult, given that multivalents normally orientate linearly so that multivalents will similarly be killed towards this mode. However 'parallel', 'indifferent' and the more stable 'convergent' modes of chain orientation were also all observed indicating that survival of some translocations in this species may be possible.

4.7 It is suggested that semi-sterile males would prove more effective than releases of completely sterile males for reducing wild populations of *H. prolixus*, because of the delayed effects of sterilizing radiation which is consequent upon the holokinetic structure of tritamine chromosomes.

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Giemsa banding of metaphase chromosomes in triatomine bugs

CONSIDERABLE progress has been made in developing the formal genetics and cytogenetics of several insect vectors of disease¹⁻³, notably with mosquitoes, houseflies and tsetse flies in which the polytene chromosomes provide suitable material for detailed analysis of chromosome morphology. The triatomine bugs (Hemiptera, Reduviidae) are medically important as vectors of Chagas' disease in the Americas, yet cytogenetic information on these insects is meagre⁴⁻⁵. These bugs present the same problems which, until recently, limited developments in mammalian cytology in that they possess a large number (typically $2n = 22$) of small, almost indistinguishable chromosomes⁶. Further, since their chromosomes are also holokinetic⁷ (that is with non-localised centromeres) they do not show any primary constrictions and it is correspondingly difficult to recognise arms of a chromosome which are readily seen in chromosomes of organisms possessing discrete centromeres. This difficulty with triatomine material has now been overcome by applying the Giemsa staining methods, which have been effectively developed by mammalian cytogenetics⁸⁻¹⁰, to embryonic cells of bugs with metaphase configuration. The technique I describe here makes possible the identification of individual chromosomes within the complements of different species of triatomine bug.

The procedure is as follows: 5-7-d-old embryos are dissected out from bug eggs, placed in hypotonic sodium citrate solution (1% for 8 min) and the cells dispersed in 0.25% trypsin in

0.02% versene. The suspension is washed in insect ringer, centrifuged and fixed in alcohol-acetic acid. Slides are prepared by air drying in the normal way¹¹. Dry slides are kept for 1 week at 20° C before heating at 60° C for 1 h in 2 × SSC (0.3 M sodium chloride; 0.03 M trisodium citrate). After rinsing in absolute ethanol, preparations are stained in 2% Giemsa (Guars 'improved' R 66) at pH 6.8 for 30 min, dried and mounted. This ASG (acetic-saline-Giemsa) technique¹² has proved to be a more reliable method for the production of G bands in bug chromosomes than other methods which have also proved effective with mammalian cells.

Figure 1 shows banding and differentiation of metaphase chromosomes in *Triatoma infestans*, and indicates the results obtainable with this technique. Similar preparations have been obtained for other *Triatoma* species and also *Rhodnius prolixus*. Preliminary analyses indicate that there are prospects for production of chromosome maps of triatomine species which could be of value in further investigations of chromosome markers and their application to problems in genetic characterisation and manipulation in the vectors of Chagas' disease.

I thank R. C. Page for assistance, H. M. Matsuda and C. Constantinos for supplying material, Professor D. S. Bernstein for his encouragement, and a grant from the Ministry of Overseas Development.

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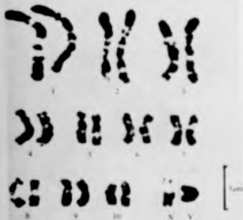


Fig. 1 Metaphase neuroblast from a *Triatoma infestans* male embryo 8 d old ($2n = XY$).

Received September 6, 1974.

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